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DIALOG(R)File 345:Inpadoc/Fam.& Legal Stat
(c) 2004 EPO. All rts. reserv.
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DIALOG(R) File 345: Inpadoc/Fam. & Legal Stat
(c) 2004 EPO. All rts. reserv.
13844934
Basic Patent (No, Kind, Date): EP 309297 A2 19890329 <No. of Patents: 119>
PATENT FAMILY:
AUSTRIA (AT)
    Patent (No, Kind, Date): AT 100465 E
                                                                                        19940215
        ANALOGE VON
                                     BRADYKININ, DESSEN SYNTHESE UND DESSEN BENUTZUNG IN DER
            THERAPIE. (German)
        Patent Assignee: UNIV TULANE (US)
        Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE; TAYLOR JOHN E
                                                                                                  A 19890328; US 173311 A
                            (No,Kind,Date): EP 89303065
            19880325
        Applic (No,Kind,Date): EP 89303065 A Addnl Info: 00334685 19940119
                                                                                                19890328
        IPC: * C07K-007/00; A61K-037/02
CA Abstract No: * 112(17)158978R; 112(19)179890W
Derwent WPI ACC No: * C 89-280003; C 89-309505
        Language of Document: English
    Patent (No, Kind, Date): AT 113961 E
                                                                                        19941115
       THERAPEUTISCHE PEPTIDE. (German)
Patent Assignee: UNIV TULANE (US)
Author (Inventor): COY DAVID H (U
        Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US) Priority (No,Kind,Date): US 100571 A 19870924
        Applic (No, Kind, Date): EP 88308916 A
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        IPC: * C07K-007/00; A61K-037/02; C07K-007/02
        CA Abstract No: * 111(11)097733N
        Derwent WPI Acc No: * C 89-095447
        Language of Document: German
                                                                                        19960715
    Patent (No,Kind,Date):
                                                      AT 139540 E
        HEILMITTELPEPTIDE (German)
        Patent Assignee: BIOMEASURE INC (US); UNIV TULANE (US)
        Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE
            SUN HYUK
                                (US)
                             (No,Kind,Date):
                                                                    us 397169
                                                                                                           19890821; US 502438 A
        Priority
            19900330
       Applic (No,Kind,Date): EP 90913117 A Addnl Info: 00489089 19960619 IPC: * C07K-007/02; C07K-007/06
                                                                                                19900817
        IPC: * C07K-007/02; C07K-007/06
CA Abstract No: * 113(19)172755T; 115(15)150377K
        Derwent WPI Acc No: * C 90-147822; C 91-087241
        Language of Document: German
                                                                                        19961015
    Patent (No.Kind,Date): AT 143372 E
        SUBSTANCÉ P ANTAGONISTE (German)
       Patent Assignee: UNIV TULANE (US)
Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US)
Priority (No,Kind,Date): US 394727 A 19890816
       Applic (No, Kind, Date): US 394727 A Addnl Info: 00438566 19960925 IPC: * C07K-007/02: C07K-007/
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        IPC: * C07K-007/02; C07K-007/22; A61K-038/08
CA Abstract No: * 115(15)151906U; 123(21)286737A
        Derwent WPI Acc No: * C 91-087240; C 95-169633
        Language of Document: German
                                                                                        19980515
    Patent (No, Kind, Date): AT 165836
        PEPTIDE ALS ARZNEIMITTEL (German)
                                              UNIV TULANE
        Patent Assignee:
             thor (Inventor): COY DAVID H
TAYLOR JOHN E (US); KIM SUN HYUK
                                                                                        (US); MOREAU JACQUES-PIERRE (US);
                                                                                        (US)
                            (No, Kind, Date):
                                                                   US 257998
                                                                                                           19881014; US 282328
        Priority
             19881209; US 317941 A
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Addnl Info: 00438519 19980506
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    123(21)286737A; 128(18)213739W
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                                 * C 89-309505; C 90-147822; C 91-087241; C
       95-169633
    Language of Document: German
AUSTRALIA (AU)
  Patent (No, Kind, Date): AU 8827102 A1 19890418
    THERAPEUTIC PEPTIDES (English)
    Patent Assignee: UNIV TÜLANE
Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE
    Priority (No, Kind, Date): WO 88US3286 A
                                                     19880923; US 100571 A
       19870924
    Applic (No, Kind, Date): AU 8827102 A
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    IPC: * C07K-007/02; C07K-007/06; C07K-007/08
Derwent WPI ACC No: * C 89-095447
    Language of Document: English
  Patent (No, Kind, Date): AU 8934146 A1 19891016
    THERAPEUTIC PEPTIDES (English)
    Patent Assignee: UNIV TULANE
    Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE; TAYLOR JOHN E;
      KIM SUN HYUK
    Priority (No, Kind, Date): WO 89US1259 A
                                                       19890327; US 173311 A
      19880325; US 282328 A
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    Applic (No, Kind, Date): AU 8934146 A
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             C07K-007/18
    Language of Document: English
  Patent (No, Kind, Date): AU 8934280 A1 19891016
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    Patent Assignee: UNIV TULANE
    Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE; TAYLOR JOHN E
    Priority (No, Kind, Date): WO 89US1216 A
                                                     19890322; US 173311 A
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    Applic (No, Kind, Date): AU 8934280 A
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    IPC: *
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    Language of Document:
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  Patent (No, Kind, Date): AU 8944949 A1 19900501
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    Patent Assignee: UNIV TULANE
    Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE; TAYLOR JOHN E;
      KIM SUN HYUK
    Priority (No, Kind, Date): WO 89US4616 A
                                                      19891013; US 257998
   19881014; US 282328 A 19881209; US 317941 A 19890707; US 397169 A 19890821 Applic (No,Kind,Date): AU 8944949 A 19891013
                                                                 19890302; us 376555
    IPC: * C07K-007/02; C07K-007/06; C07K-007/08; C07K-007/10; C07K-007/30 Derwent WPI ACC No: * C 89-309505
    Language of Document: English
  Patent (No,Kind,Date): AU 9061231 A1 19910228
    TWO-SIDED PLAYING PIECE GAME SET (English)
    Patent Assignee: LAMLE STEWART M
    Priority (No, Kind, Date): US 398172
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   Applic (No, Kind, Date): AU 9061231 A IPC: * A63F-009/20; A63F-001/02 Language of Document: English
                                                   19900822
 Patent (No, Kind, Date): AU 9062940 A1 19910403
    THERAPEUTIC PEPTIDES (English)
    Patent Assignee: BIOMEASURE INC
    Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE: KIM SUN HYUK
    Priority (No, Kind, Date): WO 90ÚS4646 A
                                                      19900817; US 397169 A
   19890821; US 502438 A 19900330
Applic (No,Kind,Date): AU 9062940 A 19900817
IPC: * C07K-005/02; C07K-005/06; C07K-005/08; C07K-005/10; C07K-007/02; C07K-007/06; C07K-007/06; C07K-007/10; C07K-007/30
CA Abstract No: * 113(19)172755T
   Derwent WPI Acc No: * C 90-147822
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Language of Document: English Patent (No,Kind,Date): AU 9514808 A1 19960926
    BOMBESIN ANALOGS (English)
   Patent Assignee: BIOMEASURE INC
   Author (Inventor): KIM SUN HYUK; MOREAU JACQUES-PIERRE
   Priority (No, Kind, Date): AU 9514808 A
                                                                           19950313; US 337127 A
       19941110
   Applic (No, Kind, Date): AU 9514808 A IPC: * C07K-007/02; A61K-038/08 CA Abstract No: * 128(18)213739W
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   Derwent WPI Acc No: * C 96-455920; C 98-229235; C 99-189718; C
       96-455920
Language of Document: English
Patent (No, Kind, Date): AU 622123 B2 19920402
   THERAPEUTIC PEPTIDES (English)
   Patent Assignee: UNIV TULANE
Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE
Priority (No,Kind,Date): WO 88US3286 A 19880923; US
                                                                             19880923; US 100571 A
       19870924
   Applic (No, Kind, Date): AU 8827102 A
                A61K-037/02; C07K-007/06; C07K-007/08
   CA Abstract No: *
                                    111(11)097733N
Derwent WPI Acc No: * C 89-095447
Language of Document: English
Patent (No, Kind, Date): AU 624566 B2 19920611
TWO-SIDED PLAYING PIECE GAME SET (English)
Patent Assignee: LAMLE STEWART M
   Author (Inventor): LAMLE STEWART M
   Priority (No, Kind, Date): US 398172 A
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   Applic (No, Kind, Date): AU 9061231 A
IPC: * G06F-015/44; A63F-009/20: A63F
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   IPC: * G06F-015/44; A63F-009/20; A63F-001/02

Derwent WPI Acc No: * G 91-059703

Language of Document: English
                                                            B2 19930701
Patent (No, Kind, Date): AU 638423
   THERAPEUTIC PEPTIDES (English)
   Patent Assignee: UNIV TÜLANE
   Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE; TAYLOR JOHN E;
       KIM SUN HYUK
   Priority (No, Kind, Date): WO 89US4616 A
                                                                               19891013; US 257998
  Priority (No,Kind,Date): WO 89US4616 A 19891013; US 257998 A 19881014; US 282328 A 19881209; US 317941 A 19890302; US 376555 A 19890707; US 397169 A 19890821

Applic (No,Kind,Date): AU 8944949 A 19891013

IPC: * C07K-007/02; C07K-007/06; C07K-007/08; C07K-007/10; C07K-007/30

CA Abstract No: * 112(17)158978R; 113(19)172755T; 115(15)150377K

Derwent WPI ACC No: * C 89-309505; C 90-147822; C 91-087241

Language of Document: English

Stent (No,Kind,Date): AU 648037 R2 19940414
                                                             B2 19940414
Patent (No,Kind,Date): AU 648037
   THERAPEUTIC PEPTIDES (English)
   Patent Assignee: BIOMEASURE INC
   Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE; KIM SUN HYUK Priority (No,Kind,Date): WO 90US4646 A 19900817; US 397169 A 19890821; US 502438 A 19900330 Applic (No,Kind,Date): AU 9062940 A 19900817
   IPC: * C07K-005/02; C07K-005/06; C07K-005/08; C07K-005/10; C07K-007/02; C07K-007/08; C07K-007/10; C07K-007/30; A61K-037/02 CA Abstract No: * 113(19)172755T; 115(15)150377K

Derwent WPI Acc No: * C 90-147822; C 91-087241

Language of Document: English
Patent (No, Kind, Date): AU 703865 B2 19990401 BOMBESIN ANALOGS (English) Patent Assignee: BIOMEASURE INC
   Author (Inventor): KIM SUN HYUK; MOREAU JACQUES-PIERRE
   Priority (No, Kind, Date): AU 9514808 A
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   Applic (No,Kind,Date): AU 9514808 A IPC: * C07K-007/02; A61K-038/08 Derwent WPI Acc No: * C 96-455920
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   Language of Document: English
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CANADA (CA)
     Patent (No, Kind, Date): CA 2008454 AA 19900902
        THERAPEUTIC PEPTIDES (English; French)
        Patent Assignee: UNIV TULANE (Author (Inventor): COY DAVID H
                                                     (US)
                                                      (US); MOREAU JACQUES P (US); TAYLOR
       JOHN E (US); KIM SUN H (US)
Priority (No,Kind,Date): US 317941 A 1989
19890707; US 397169 A 19890821
Applic (No,Kind,Date): CA 2008454 A 19900
National Class: * D3530000706 M; 16701038 S
                                                                   19890302; US 376555 A
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        IPC: * C07K-007/06; A61K-037/02
        CA Abstract No: * 113(19)172755T
    Derwent WPI Acc No: * C 90-147822
Language of Document: English
Patent (No, Kind, Date): CA 2023460 AA 19910224
TWO-SIDED PLAYING PIECE GAME SET (English; French)
       Patent Assignee: LAMLE STEWART M (US)
Author (Inventor): LAMLE STEWART M (US)
Priority (No,Kind,Date): US 398172 A
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       Applic (No, Kind, Date): CA 2023460 A National Class: * D42720065 M
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       IPC: * A63F-009/20
       Language of Document:
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CA 2039175 AA
    Patent (No, Kind, Date):
       THERAPEUTIC PEPTIDES (English; French)
       Patent Assignee: UNIV TÜLANE (US)
       Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE
       Priority (No, Kind, Date): US 394727 A
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      Applic (No, Kind, Date): CA 2039175 A 19900816
National Class: * D2530000708 M; 530000506 S; 530000502 S; 530000508 S
IPC: * C07K-007/08; C07K-007/06; C07K-007/02; C07K-005/00
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      CA Abstract No: * 115(15)1519060
   Derwent WPI ACC No: * C 91-087240
Language of Document: English
Patent (No,Kind,Date): CA 2064896 AA
                                                             19910222
      THERAPEUTIC PEPTIDES (English; French)
      Patent Assignee: BIOMEASURE INC (US); UNIV TULANE (US)
      Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE
                                                                                               (US); KIM
      SUN H (US)
Priority (No,Kind,Date): US 397169 A
                                                                 19890821; US 502438
         19900330
      Applic (No, Kind, Date): CA 2064896 A 19900817

IPC: * C07K-007/06; C07K-007/00; C07K-005/00; C07K-007/30

CA Abstract No: * 113(19)172755Τ; 115(15)150377Κ
   Derwent WPI ACC No: * C 90-147822; C 91-087241
Language of Document: English
Patent (No, Kind, Date): CA 1335622 A1 19950516
      BRADYKININ ANALOGS CONTAINING A NON-PEPTIDE BOND (English; French)
      Patent Assignee: UNIV TULANE (US)
      Author (Inventor):
                                  COY DAVID H (US); MOREAU JACQUES-PIERRE (US);
         TAYLOR JOHN E
                             (US)
      Priority (No, Kind, Date): US 173311 A
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     Applic (No, Kind, Date):
                                       CA 594845 A
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     National Class: * D1530000718 M; 167010346 S
     IPC: * C07K-007/18; A61K-037/42; A61K-037/43
CA Abstract No: * 112(17)158978R; 112(19)179890W; 123(21)286737A;
        128(18)213739w; 129(02)016394z
     Derwent WPI Acc No: * C 89-280003; C 98-229235; C 98-296827; C 99-189718
                                                               89-309505; C 95-169633; C
     Language of Document: English
CZECH REPUBLIC (CZ)
  Patent (No, Kind, Date): CZ 9004028 A3
     LINEAR PEPTIDE (Czech; English)
Patent Assignee: UNIV TULANE
                                                  (US)
     Author (Inventor): COY DAVID H
                                                    (US); MOREAU JACQUES-PIERRE (US)
     Priority (No, Kind, Date): US 394727 A
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Applic (No, Kind, Date): CZ 904028 A
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Derwent WPI Acc No: * C 91-087240; C 95-169633
Language of Document: Czech; Slovak
    Patent (No, Kind, Date): CZ 285319 B6 19990714
      LINEAR PEPTIDE (Czech; English)
Patent Assignee: UNIV TULANE (US)
Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US)
Priority (No,Kind,Date): US 394727 A 19890816
Applic (No,Kind,Date): CZ 904028 A 19900816
IPC: * CO7K-007/02; CO7K-007/22; A61K-038/08
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       Derwent WPI Acc No: * C 91-087240; C 95-169633
Language of Document: Czech; Slovak
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Patent Assignee: UNIV TULANE (US)
Author (Inventor): COY DAVID H (US);
Priority (No, Kind, Date): US 394727 A
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      IPC: * C07K-007/02; C07K-007/22
CA Abstract No: * 115(15)151906U; 123(21)286737A
Derwent WPI Acc No: * C 91-087240; C 95-169633
Language of Document: Czech; Slovak
GERMAN DEMOCRATIC REPUBLIC (DD)
Patent (No, Kind, Date): DD 298411 A5 19920220
       THERAPEUTISCHE PEPTIDE (German)
       Patent Assignee: UNIV TULANE (US)
       Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US)
       Priority (No, Kind, Date): US 394727 A
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      Applic (No, Kind, Date): DD 343501 A 199008

IPC: * C07K-007/06; A61K-037/02; C07K-007/02

CA Abstract No: * 115(15)151906U

Derwent WPI Acc No: * C 91-087240

Language of Document: German
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GERMANY (DE)
    Patent (No, Kind, Date): DE 3852086 CO 19941215
      THERAPEUTISCHE PEPTIDE. (German)
Patent Assignee: UNIV TULANE (US)
Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US)
Priority (No,Kind,Date): US 100571 A 19870924
Applic (No,Kind,Date): DE 3852086 A 19880926
IPC: * CO7K-007/00; A61K-037/02; C07K-007/02
CA Abstract No: * 111(11)097733N
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       Derwent WPI Acc No: * C 89-095447
Language of Document: German
    Patent (No, Kind, Date): DE 68912376 CO 19940303
       ANALOGE VON BRADYKININ, DESSEN SYNTHESE UND DESSEN BENUTZUNG IN DER
          THERAPIE: (German)
       Patent Assignee: UNIV TULANE (US)
       Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US); TAYLOR JOHN E (US)
       Priority (No, Kind, Date): US 173311 A
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       Applic (No,Kind,Date): EP 89303065 A IPC: * C07K-007/00; A61K-037/02
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       CA Abstract No: * 112(17)158978R; 112(19)179890W
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Language of Document: German
Patent (No, Kind, Date): DE 68928667 CO 19980610
       PEPTIDE ALS ARZNEIMITTEL (German)
       Patent Assignee: UNIV TULANE (US)
       Author (Inventor): COY DAVID (US); MOREAU JACQUES-PIERRE (US);
       TAYLOR JOHN (US); KIM SUN (US)
Priority (No, Kind, Date): US 257998 A 19881014; US 282328 A 19881209; US 317941 A 19890302; US 376555 A 19890707; US 397169
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19890821; wo 89us4616 w
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      Applic (No, Kind, Date): DE 68928667 A 19891013
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CA Abstract No: * 112(17)158978R; 113(19)172755T; 115(15)150377K;
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Derwent WPI Acc No: * C 89-309505; C 90-147822; C 91-087241; C 95-169633; C 98-229235

Language of Document: German
  Patent (No, Kind, Date): DE 69027533 CO 19960725
      HEILMITTELPEPTIDE (German)
      Patent Assignee: BIOMEASURE INC (US); UNIV TULANE (US)
      Author (Inventor): COY DAVID (US); MOREAU JACQUES-PIERRE (US); KIM
           SUN
                     (US)
     Priority (No,Kind,Date): US 397169
19900330; WO 90US4646 W 1990081
Applic (No,Kind,Date): DE 69027533
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      IPC: * C07K-007/02; C07K-007/06
CA Abstract No: * 113(19)172755T;
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      Derwent WPI Acc No: * C 90-147822; C 91-087241
      Language of Document: German
 Patent (No, Kind, Date): DE 69028692 CO 19961031
      SUBSTANCE P ANTAGONISTE (German)
     Patent Assignee: UNIV TULANE (US)
Author (Inventor): COY DAVID (US); MOREAU JACQUES-PIERRE (US)
Priority (No, Kind, Date): US 394727 A 19890816; WO 90US4633
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     Applic (No,Kind,Date): DE 69028692
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                       C07K-007/02; C07K-007/22; A61K-038/08
     CA Abstract No: * 115(15)151906Ú; 123(21)286737A
     Derwent WPI Acc No: * C 91-087240; C 95-169633
Language of Document: German
Patent (No, Kind, Date): DE 3852086 T2 19950518 THERAPEUTISCHE PEPTIDE. (German)
Patent Assignee: UNIV TULANE (US)
     Author (Inventor): COY DAVID H
                                                                                 (US); MOREAU JACQUES-PIERRE (US)
     Priority (No, Kind, Date): US 100571 A
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    Applic (No, Kind, Date): DE 3852086 A 198809

IPC: * C07K-007/00; A61K-038/00; C07K-007/02

CA Abstract No: * 111(11)097733N

Derwent WPI Acc No: * C 89-095447

Language of Document: German
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Patent (No, Kind, Date): DE 68912376 T2 19940707
     ANALOGE VON BRADYKININ, DESSEN SYNTHESE UND DESSEN BENUTZUNG IN DER
         THERAPIE. (German)
     Patent Assignee: UNIV TULANE (US)
     Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US); TAYLOR JOHN E (US)
     Priority (No, Kind, Date): US 173311 A
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    Applic (No, Kind, Date): DE 68912376 IPC: * C07K-007/00: A61K-037/02
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    IPC: * C07K-007/00; A61K-037/02
CA Abstract No: * 112(17)158978R; 112(19)179890W
Derwent WPI Acc No: * C 89-280003; C 89-309505
Language of Document: German
Patent (No, Kind, Date): DE 68928667 T2 19981001
     PEPTIDE ALS ARZNEIMITTEL (German)
     Patent Assignee: UNIV TULANE
                                                                              (US)
     Author (Inventor): COY DAVID (US); MOREAU JACQUES-PIERRE (US);
    TAYLOR JOHN (US); KIM SUN (US)
Priority (No, Kind, Date): US 257998 A
19881209; US 317941 A 19890302; US
                                                                                                     19881014; US 282328
                                                                   19890302; US 376555 A
                                                                                                                               19890707; us 397169
   A 19890821; WO 89US4616 W 19891013

Applic (No,Kind,Date): DE 68928667 A 19891013

IPC: * CO7K-007/02; CO7K-014/595; A61K-038/16

CA Abstract No: * 112(17)158978R; 113(19)172755T; 115(15)150377K; 123(21)286737A; 128(18)213739W; 129(02)016394Z

Derwent WPI Acc No: * C 89-309505; C 90-147822; C 91-087241; C 95-169633: C 98-220235; C 98-220235;
                                                                                               90-147822; C 91-087241; C
         95-169633; c 98-229235; c 98-296827
    Language of Document: German
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Patent (No, Kind, Date): DE 69027533 T2 19961219
     HEILMITTELPEPTIDE (German)
     Patent Assignee: BIOMEASURE INC (US); UNIV TULANE (US)
     Author (Inventor): COY DAVID (US); MOREAU JACQUES-PIERRE (US); KIM
             (US)
     Priority (No, Kind, Date): US 397169 A 19900330; WO 90US4646 W 19900817
                                                            19890821; US 502438 A
     Applic (No,Kind,Date): DE 69027533 A IPC: * C07K-007/02; C07K-007/06
                                                            19900817
     CA Abstract No: * 113(19)172755T; 115(15)150377K
     Derwent WPI Acc No: * C 90-147822; C 91-087241
     Language of Document: German
   Patent (No, Kind, Date): DE 69028692 T2 19970220
     SUBSTANCE P ANTAGONISTE (German)
     Patent Assignee: UNIV TULANE (US)
Author (Inventor): COY DAVID (US); MOREAU JACQUES-PIERRE
Priority (No,Kind,Date): US 394727 A 19890816; WO 90US46
                                                          19890816; wo 90US4633
        19900816
     Applic (No, Kind, Date): DE 69028692
                                                            19900816
     IPC: * C07K-007/02; C07K-007/22; A61K-038/08
CA Abstract No: * 115(15)151906U; 123(21)286737A
     Derwent WPI Acc No: * C 91-087240; C 95-169633
Language of Document: German
DENMARK (DK)
  Patent (No,Kind,Date): DK 9100663 A 1
TERAPEUTISK VIRKSOMME PEPTIDER (Danish)
                                                        19910614
     Patent Assignee: UNIV TULANE (US)
     Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE; TAYLOR JOHN E;
        KIM SUN HYUK
     Priority (No, Kind, Date): US 257998 A 19881014; 19881209; US 317941 A 19890302; US 376555 A
                                                            19881014; US 282328
                                                                          19890707; US 397169
     A 19890821; WO 89US4616 A 19891013
Applic (No,Kind,Date): DK 91663 A 199
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     IPC: * CO7K-007/30
CA Abstract No: * 112(17)158978R; 113(19)172755T
     Derwent WPI Acc No: * C 89-309505; C 90-147822; C 91-087241
  Language of Document: Danish
Patent (No,Kind,Date): DK 8902494 A 19890720
THERAPEUTISK VIRKSOMME PEPTIDER, ISAER BOMBESINANTAGONISTISKE OG
       LITORINANTAGONISTISKE PEPTIDER SAMT FREMGANGSMAADE TIL FREMSTILLING
        DERAF (Danish)
                            UNIV TULANE (US)
     Patent Assignee:
     Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE
     Priority (No,Kind,Date): US 100571 A 19880923
                                                          19870924; WO 88US3286 A
     Applic (No, Kind, Date): DK 892494 A IPC: * C07K-007/08
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     CA Abstract No: * 111(11)097733N
  Derwent WPI Acc No: * C 89-095447
Language of Document: Danish
Patent (No,Kind,Date): DK 9100663 A0
     TERAPEUTÍSK VÍRKSOMME PEPTIDER (Danish)
     Patent Assignee: UNIV TULANE (US)
     Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE; TAYLOR JOHN E;
        KIM SUN HYUK
     Priority (No, Kind, Date): US 257998 A 19881014; 19881209; US 317941 A 19890302; US 376555 A
                                                            19881014; US 282328
                                                                         19890707; US 397169
     A 19890821; WO 89US4616 A 19891013
Applic (No, Kind, Date): DK 91663 A 19910412
IPC: * CO7K-007/30
CA Abstract No: * 112(17)158978R; 113(19)172755T
     Derwent WPI Acc No: * C 89-309505; C 90-147822; C 91-087241 Language of Document: Danish
  Patent (No, Kind, Date): DK 8902494 AO 19890523
THERAPEUTISK VIRKSOMME PEPTIDER, ISAER BOMBESINANTAGONISTISKE OG
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        DERAF (Danish)
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Patent Assignee: UNIV TULANE (US)
     Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE
     Priority (No,Kind,Date): US 100571 A 19880923
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     Applic (No,Kind,Date): DK 892494 A IPC: * C07K-007/08
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     Derwent WPI Acc No: *
                                C 89-095447
     Language of Document: Danish
  Patent (No, Kind, Date): DK 438566 T3
                                                    19961111
     SUBSTANS P-ANTAGONISTER (Danish)
     Patent Assignee: UNIV TULANE (US)
Author (Inventor): COY DAVID H (U
                                              (US); MOREAU JACQUES-PIERRE (US)
     Priority (No, Kind, Date): US 394727
                                                        19890816
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     Applic (No, Kind, Date): DK 9090912128 A 19900816

IPC: * C07K-005/02; C07K-005/06; C07K-005/08; C07K-005/10; C07K-007/02; C07K-007/08

CA Abstract No: * 115(15)151906U; 123(21)286737A

Derwent WPT Acc No: * 6 91-087340; C 05 160633
     Derwent WPI Acc No: * C 91-087240; C 95-169633
     Language of Document: Danish
  Patent (No,Kind,Date): DK 489089 T3 19960729
     TERAPEUTISKE PEPTIDER (Danish)
     Patent Assignee: UNIV TULANE (US); BIOMEASURE INC (US)
     Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US); KIM
       SUN HYUK (US)
     Priority (No, Kind, Date): US 397169 A
                                                        19890821; US 502438
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     Applic (No,Kind,Date): DK 9090913117 A
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               C07K-005/02; C07K-005/06; C07K-005/08; C07K-005/10; C07K-007/02
    ; C07K-007/06; C07K-007/08; C07K-007/10; C07K-007/30
CA Abstract No: * 113(19)172755T; 115(15)150377K; 128(18)213739W
Derwent WPI Acc No: * C 90-147822; C 91-087241; C 98-229235
     Language of Document: Danish
EUROPEAN PATENT OFFICE (EP)
  Patent (No, Kind, Date): EP 414512 A1 19910227
     TWO-SIDED PLAYING PIECE GAME SET (English; French; German)
     Patent Assignee: LAMLE STEWART M (US)
    Author (Inventor): LAMLE STEWART M
    Priority (No,Kind,Date): US 398172 A Applic (No,Kind,Date): EP 90309187 A
                                                        19890823
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    Designated States: (National) AT; BE; CH; DE; DK; ES; FR; GB; GR; IT;
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  Derwent WPI ACC No:; G 91-059703
Language of Document: English
Patent (No,Kind,Date): EP 438519 A1 19910731
    THERAPEUTIC PEPTIDES (English; French; German)
    Patent Assignee: UNIV TÜLANE
                                           (US)
    Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US);
    TAYLOR JOHN E (US); KIM SUN HYUK (US) Priority (No, Kind, Date): WO 89US4616 W
                                                  (US)
                                                           19891013; US 257998
       19881014; US 282328 A 19
A 19890707; US 397169 A
                                     19881209; US 317941 A
                                                                      19890302; us 376555
                                           19890821
    Applic (No, Kind, Date): EP 89912292 A
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    Designated States: (National) AT; BE; CH; DE; FR; GB; IT; LI; LU; NL;
    IPC: * C07K-007/02; C07K-007/06; C07K-007/08; C07K-007/10; C07K-007/30 CA Abstract No: * 112(17)158978R; 113(19)172755T
    Derwent WPI Acc No: * C 89-309505; C 90-147822; C 91-087241
    Language of Document: English
  Patent (No, Kind, Date): EP 438566 A1 19910731
    THERAPEUTIC PEPTIDES (English; French; German)
    Patent Assignee: UNIV TULANE (US)
Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE
Priority (No,Kind,Date): WO 90US4633 W 19900816; US 39472
                                                          19900816; US 394727 A
       19890816
    Applic (No, Kind, Date): EP 90912128 A
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    Designated States: (National) AT; BE; CH; DE; DK; ES; FR; GB; IT; LI;
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LU; NL; SE IPC: * C07K-005/02; C07K-005/06; C07K-005/08; C07K-005/10; C07K-007/02
      ; C07K-007/06; C07K-007/08
   Derwent WPI Acc No: * C 91-087240
Language of Document: English
Patent (No, Kind, Date): EP 489089 A1 19920610 THERAPEUTIC PEPTIDES (English; French; German)
   Patent Assignee: BIOMEASURE INC (US); UNIV TULANE (US)
   Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US); KIM
     SUN HYUK
                  (US)
   Priority (No, Kind, Date): WO 90US4646 W
                                                          19900817; US 397169 A
     19890821; US 502438 A
                                     19900330
   Applic (No, Kind, Date): EP 90913117
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   Designated States: (National) AT; BE; CH; DE; DK; ES; FR; GB; IT; LI;
  LU; NL; SE

IPC: * C07K-005/02; C07K-005/06; C07K-005/08; C07K-005/10; C07K-007/02; C07K-007/08; C07K-007/10; C07K-007/30

CA Abstract No: * 113(19)172755T; 115(15)150377K

Derwent WPI Acc No: * C 90-147822; C 91-087241

Language of Document: English
Patent (No, Kind, Date): EP 309297 A2 19890329
   THERAPEUTIC PEPTIDES (English; French; German)
  Patent Assignee: UNIV TULANE (US)
Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE
Priority (No,Kind,Date): US 100571 A 19870924
Applic (No,Kind,Date): EP 88308916 A 19880926
  Designated States: (National) AT; BE; CH; DE; ES; FR; GB; GR; IT; LI;
  LU; NL; SE
IPC: * CO7K-
            C07K-007/00; A61K-037/02
   CA Abstract No: ; 111(11)097733N
Derwent WPI Acc No: ; C 89-095447
Language of Document: English
Patent (No,Kind,Date): EP 334685 A2 19890927
   BRADYKININ ANALOGUES, THEIR SYNTHESIS AND THEIR USE IN THERAPY (English
      French; German)
  Patent Assignee: UNIV TULANE (US)
  Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE; TAYLOR JOHN E
  Priority (No, Kind, Date): US 173311 A
                                                       19880325
  Applic (No, Kind, Date): EP 89303065
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  Designated States: (National) AT; BE; CH; DE; ES; FR; GB; GR; IT; LI;
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IPC: * C07K-007/00; A61K-037/02
  CA Abstract No: ; 112(19)179890W
  Derwent WPI Acc No: ; C 89-280003
Language of Document: English
Patent (No, Kind, Date): EP 309297 A3
                                                 19900704
  THERAPEUTIC PEPTIDES (English; French; German)
  Patent Assignee: UNIV TULANE (US)
Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE
  Priority (No,Kind,Date): US 100571 A Applic (No,Kind,Date): EP 88308916 A
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  Designated States: (National) AT; BE; CH; DE; ES; FR; GB; GR; IT; LI;
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IPC: * C07K
            C07K-007/00; A61K-037/02; C07K-007/02
  CA Abstract No: * 111(11)097733N
Derwent WPI ACC No: * C 89-095447
Language of Document: English
Patent (No,Kind,Date): EP 334685 A3
                              EP 334685 A3 19910130
  BRADYKINÍN ANÁLOGUES, THEIR SYNTHESIS AND THEIR USE IN THERAPY (English
     ; French; German)
  Patent Assignee: UNIV TULANE (US)
  Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE; TAYLOR JOHN E
  Priority (No, Kind, Date): US 173311 A
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  Applic (No, Kind, Date): EP 89303065 A
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IPC: * CO7K
           C07K-007/00; A61K-037/02
  CA Abstract No: * 112(17)158978R; 112(19)179890w
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Derwent WPI Acc No: * C 89-280003; C 89-309505
   Language of Document: English
Patent (No, Kind, Date): EP 438519 A4 19911030
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   Patent Assignee:
                      UNIV TULANE
                                     (US)
  Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US);
     TAYLOR JOHN E (US); KIM SUN HYUK
  Priority (No, Kind, Date): WO 89US4616 W
                                                   19891013; US 257998
     19881014; US 282328 A
                                19881209; US 317941 A
                                                              19890302; us 376555
         19890707; US 397169 A
                                      19890821
  Applic (No, Kind, Date): EP 89912292 A
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  Designated States: (National) AT; BE; CH; DE; FR; GB; IT; LI; LU; NL;
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  IPC: * C07K-007/02; C07K-007/06; C07K-007/08; C07K-007/10; C07K-007/30 CA Abstract No: * 112(17)158978R; 113(19)172755T
  Derwent WPI Acc No: * C 89-309505; C 90-147822; C 91-087241
  Language of Document: English
Patent (No, Kind, Date):
                           EP 438566 A4 19930331
  THERAPEUTIC PEPTIDES (English; French; German)
  Patent Assignee: UNIV TULANE
                                     (US)
  Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US)
  Priority (No, Kind, Date): WO 90US4633 W
                                                   19900816; US 394727
     19890816
  Applic (No,Kind,Date): EP 90912128 A
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  Designated States: (National) AT; BE; CH; DE; DK; ES; FR; GB; IT; LI;
           C07K-005/02; C07K-005/06; C07K-005/08; C07K-005/10; C07K-007/02
  ; C07K-007/06; C07K-007/08
CA Abstract No: * 115(15)151906U
  Derwent WPI Acc No: * C 91-087240
Language of Document: English
Patent (No, Kind, Date):
                           EP 489089 A4 19920624
  THERAPEUTIC PEPTIDES (English; French; German)
  Patent Assignee: BIOMEASURE INC (US); UNIV TULANE (US)
  Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US); KIM
    SUN HYUK (US)
  Priority (No,Kind,Date): WO 90US4646 W 19890821; US 502438 A 19900330 Applic (No,Kind,Date): EP 90913117 A
                                                   19900817; US 397169 A
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  Designated States: (National) AT; BE; CH; DE; DK; ES; FR; GB; IT; LI;
  IPC: * C07K-005/02; C07K-005/06; C07K-005/08; C07K-005/10; C07K-007/02; C07K-007/06; C07K-007/08; C07K-007/10; C07K-007/30 CA Abstract No: * 113(19)172755T; 115(15)150377K
  Derwent WPI Acc No: * C 90-147822; C 91-087241
Language of Document: English
Letent (No,Kind,Date): EP 309297 B1 19941109
Patent (No, Kind, Date):
  THERAPEUTIC PEPTIDES. (English; French; German)
  Patent Assignee: UNIV TULANE
                                    (US)
  Author (Inventor): COY DAVID H
                                       (US); MOREAU JACQUES-PIERRE (US)
                               US 100571 A
  Priority (No, Kind, Date):
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  Applic (No, Kind, Date):
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                                                19880926
  Designated States: (National) AT; BE; CH; DE; ES; FR; GB; GR; IT; LI;
  LU; NL; SE
IPC: * C07K-007/00; A61K-037/02; C07K-007/02
  CA Abstract No: *
                       111(11)097733N; 123(21)286737A; 128(18)213739W;
    129(02)016394Z
  Derwent WPI Acc No: * C 89-095447; C 95-169633; C 98-229235; C
    98-296827; C 99-189718
  Language of Document: English
Patent (No, Kind, Date): EP 334685 B1 19940119
  BRADYKININ ANALOGUES, THEIR SYNTHESIS AND THEIR USE IN THERAPY (English
     French; German)
 Patent Assignee: UNIV TULANE (US)
Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US);
    TAYLOR JOHN E
                    (US)
  Priority (No, Kind, Date): US 173311 A
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 Applic (No, Kind, Date): EP 89303065 A
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Designated States: (National) AT; BE; CH; DE; ES; FR; GB; GR; IT; LI;
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IPC: * C07K-007/00; A61K-037/02
CA Abstract No: * 112(17)158978R; 112(19)179890W
    Derwent WPI Acc No: * C 89-280003; C 89-309505
    Language of Document: English
  Patent (No, Kind, Date): EP 438519 B1 19980506 THERAPEUTIC PEPTIDES (English; French; German)
    Patent Assignee: UNIV TŪLANE
    Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US);
    TAYLOR JOHN E (US); KIM SUN HYUK (U
Priority (No, Kind, Date): US 257998 A
                                                 (US)
                                                      19881014; US 282328
                                    19890302; US 376555 A
                                                                   19890707; US 397169
       19881209; US 317941 A
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            19890821; wo 89us4616 w
    Applic (No, Kind, Date): EP 89912292 A
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    Designated States: (National) AT; BE; CH; DE; FR; GB; IT; LI; LU; NL;
    IPC: * C07K-007/02; C07K-014/595; A61K-038/16
CA Abstract No: * 112(17)158978R; 113(19)172755T; 115(15)150377K;
       123(21)286737A; 128(18)213739W
    Derwent WPI Acc No: * C 89-309505; C 90-147822; C 91-087241; C
       95-169633
    Language of Document: English
  Patent (No, Kind, Date): EP 438566 B1 19960925
    SUBSTANCE P ANTAGONISTS (English; French; German)
    Patent Assignee: UNIV TULANE (US)
    Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US) Priority (No, Kind, Date): US 394727 A 19890816; WO 90US4633 W
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    Designated States: (National) AT; BE; CH; DE; DK; ES; FR; GB; IT; LI;
             C07K-007/02; C07K-007/22; A61K-038/08
    CA Abstract No: * 115(15)1519060; 123(21)286737A
    Derwent WPI Acc No: * C 91-087240; C 95-169633
Language of Document: English
Lent (No,Kind,Date): EP 489089 B1 19960619
  Patent (No, Kind, Date):
    THERAPEUTIC PEPTIDES (English; French; German)
    Patent Assignee: BIOMEASURE INC (US); UNIV TULANE
    Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US); KIM
       SUN HYUK (US)
    Priority (No,Kind,Date): WO 90US4646 W
                                                        19900817: US 502438 A
    19900330; US 397169 A 19890821
Applic (No,Kind,Date): EP 90913117
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    LU; NL; SE
IPC: * C07K-007/02; C07K-007/06
CA Abstract No: * 113(19)172755T; 115(15)150377K
    Derwent WPI Acc No: * C 90-147822; C 91-087241
    Language of Document: English
SPAIN (ES)
  Patent (No, Kind, Date): ES 2061977 T3 19941216
    COMPUESTOS ANALOGOS A BRADIQUININA, SU SINTESIS Y SU USO EN TERAPIA.
       (Spanish)
    Patent Assignee: UNIV TULANE
    Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US);
       TAYLOR JOHN E (US)
                                                      19880325
    Priority (No, Kind, Date): US 173311 A
    Applic (No, Kind, Date): ES 89303065 EP 19890328
    Addnl Info: 0334685 EP patent valid in AT IPC: * C07K-007/00; A61K-037/02 CA Abstract No: * 112(17)158978R; 112(19)179890W Derwent WPI Acc No: * C 89-280003; C 89-309505 Language of Document: Spanish
  Patent (No, Kind, Date): ES 2065336 T3 19950216
    PEPTIDOS TERAPEUTICOS. (Spanish)
    Patent Assignee: UNIV TULANE
    Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US)
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Priority (No, Kind, Date): US 100571 A
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     Applic (No, Kind, Date): ES 88308916 EP 19880926
    Addnl Info: 0309297 EP patent valid in AT IPC: * C07K-007/00; A61K-037/02; C07K-007/02 CA Abstract No: * 111(11)097733N Derwent WPI Acc No: * C 89-095447 Language of Document: Spanish
  Patent (No, Kind, Date): ES 2090140
                                            T3 19961016
     PEPTIDOS TERAPEUTICOS. (Spanish)
     Patent Assignee: BIOMEASURE INC
                                            (US); UNIV TULANE (US)
    Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE
                                                                              (US); KIM
       SUN HYUK (US)
     Priority (No, Kind, Date): US 397169 A
                                                     19890821; US 502438 A
       19900330
    Applic (No, Kind, Date): ES 90913117
                                                EP 19900817
    Addnl Info: 0489089 EP patent valid in AT
    IPC: * C07K-007/02; C07K-007/06
    CA Abstract No: * 113(19)172755T; 115(15)150377K
    Derwent WPI Acc No: * C 90-147822; C 91-087241
    Language of Document: Spanish
  Patent (No, Kind, Date): ES 2094160 T3 19970116
    PEPTIDOS TERAPEUTICOS, EN PARTICULAR ANALOGOS DEL PEPTIDO SUBSTANCIA P.
       (Spanish)
    Patent Assignee:
                         UNIV TULANE
    Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US)
    Priority (No, Kind, Date): US 394727 Á
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    Applic (No, Kind, Date): ES 90912128 EP
    Addnl Info: 0438566 EP patent valid in AT IPC: * C07K-007/02; C07K-007/22; A61K-038/08
    CA Abstract No: * 115(15)151906U; 123(21)286737A
Derwent WPI Acc No: * C 91-087240; C 95-169633
Language of Document: Spanish
FINLAND (FI)
  Patent (No, Kind, Date): FI 8902507 A
                                                 19890523
    TERAPEUTISKA PEPTIDER. (Swedish)
    Patent Assignee: UNIV TULANE (US); COY DAVID HOWARD (US); MOREAU
    JACQUES PIERRE (US)
Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE
    Priority (No,Kind,Date): US 100571 A 19880923
                                                    19870924; WO 88US3286 A
    Applic (No,Kind,Date): FI 892507 A
                                                  19890523
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    Derwent WPI Acc No: * C 89-095447
    Language of Document: Finnish; Swedish
  Patent (No, Kind, Date): FI 8902507
                                            A0 19890523
    TERAPEUTÍSKA PEPTIDER. (Swedish)
Patent Assignee: UNIV TULANE (US); COY DAVID HOWARD (US); MOREAU
    Patent Assignee:
      JACQUES PIERRE
                         (US)
    Author (Inventor):
                           COY DAVID H (US); MOREAU JACQUES-PIERRE
    Priority (No, Kind, Date): US 100571 A 19870924; WO 88US3286
      19880923
    Applic (No Kind, Date): FI 892507 A
                                                  19890523
    IPC: *
             C07K
 Derwent WPI Acc No: * C 89-095447
Language of Document: Finnish; Swedish
Patent (No,Kind,Date): FI 9004153 A0 1
                                                19900822
    DUBBELSIDIGA SPELDELAR OMFATTANDE SPEL. (Swedish)
    Patent Assignee: LAMLE STEWART MILTON
                                                  (US)
    Author (Inventor): LAMLE STEWART MILTON
    Priority (No, Kind, Date): US 398172 A
                                                    19890823
    Applic (No, Kind, Date): FI 904153 A IPC: * A63F
                                                  19900822
 Language of Document: Finnish; Swedish Patent (No, Kind, Date): FI 9101780 AO 19910412
    TERAPEUTISKA PEPTIDER. (Swedish)
    Patent Assignee: UNIV TULANE (US)
    Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US);
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TAYLOR JOHN E (US); KIM SUN HYUK (US)
     Priority (No, Kind, Date): US 257998 A
                                                              19881014; US 282328 A
                                          19890302; US 376555 A
                                                                             19890707; US 397169
        19881209; US 317941 A
                                                   19891013
             19890821; WO 89US4616 A
     Applic (No,Kind,Date): FI 911780 A IPC: * CO7K
                                                           19910412
     CA Abstract No: * 112(17)158978R; 113(19)172755T
  Derwent WPI Acc No: * C 89-309505; C 90-147822; C 91-087241 Language of Document: Finnish; Swedish Patent (No, Kind, Date): FI 9200737 A0 19920220
     TERAPEUTÍSKA PEPTIDER. (Swedish)
     Patent Assignee: BIOMEASURE INC
                                                    (US); UNIV TULANE (US)
     Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE
        SUN HYUK
                     (US)
     Priority (No,Kind,Date): US 397169 A
19900330; WO 90US4646 A 19900817
                                                       Α
                                                              19890821; US 502438
     Applic (No, Kind, Date): FI 92737 A
               C07K
  CA Abstract No: * 113(19)172755T; 115(15)150377K
Derwent WPI Acc No: * C 90-147822; C 91-087241
Language of Document: Finnish; Swedish
Patent (No, Kind, Date): FI 100719 B1 19980213
     FOERFARANDE FOER FRAMSTAELLNING AV TERAPEUTISKT ANVAENDBARA,
        MODIFIERADE BOMBESIN- OCH LITORINANTAGONISTPEPTIDER (Swedish)
     Patent Assignee: UNIV TULANE (US); COY DAVID HOWARD
                                                                                 (US); MOREAU
     JACQUES PIERRE (US)
Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US)
     Priority (No,Kind,Date): US 100571 A 19880923
                                                              19870924; WO 88US3286 W
     Applic (No, Kind, Date): FI 892507 A 1989052 IPC: * C07K-007/02; C07K-007/06; C07K-007/08
                                                           19890523
     CA Abstract No: *
                              111(11)097733N; 123(21)286737A; 128(18)213739W;
        129(02)016394Z
     Derwent WPI Acc No: * C 89-095447; C 95-169633; C 98-229235; C
        98-296827
  Language of Document: Finnish; Swedish Patent (No, Kind, Date): FI 104252 B1 19991215
     MENETELMAE TERAPEUTTISESTI KAEYTTOEKELPOISTEN PEPTIDIEN KEMIALLISEKSI
        SYNTETISOIMISEKSI KIINTEAESSAE FAASISSA FOERFARANDE FOER KEMISK
        SYNTES I FAST FAS AV TERAPEUTISKT ANVAENDBARA PEPTIDER (Swedish)
     Patent Assignee: UNIV TULANE (US)
     Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US); TAYLOR JOHN E (US); KIM SUN HYUK (US)
Priority (No, Kind, Date): US 257998 A 19881014; US 282328 A
                                         19890302; US 376555 A
                                                                             19890707; US 397169
        19881209; ÚS 317941 A
             19890821; wo 89US4616 W
                                                   19891013
     Applic (No, Kind, Date): FI 911780 A
                                                          19910412
               C07K-007/02; C07K-007/06
     CA Abstract No: * 112(17)158978R; 113(19)172755T; 115(15)150377K; 123(21)286737A; 128(18)213739W; 129(02)016394Z

Derwent WPI Acc No: * C 89-309505; C 90-147822; C 91-087241; C 95-169633; C 98-229235; C 98-296827; C 99-189718

Language of Document: Finnish; Swedish
GREECE (GR)
                                                           19911230
  Patent (No, Kind, Date): GR 90100613 A
     THERAPEUTICAL PEPTIDES (English)
     Patent Assignee: UNIV TULANE
     Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE Priority (No, Kind, Date): US 394727 A 19890816
     Applic (No, Kind, Date): GR 100613 A 19900816

IPC: * C07K-005/02; C07K-005/06; C07K-005/08; C07K-005/10; C07K-007/02; C07K-007/06; C07K-007/08; C07K-007/22

CA Abstract No: * 115(15)151906U; 123(21)286737A
     Derwent WPI Acc No: * C 91-087240; C 95-169633
     Language of Document: Greek
HONG KONG (HK)
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Patent (No, Kind, Date): HK 1010785 A1 19990625
     THERAPEUTIC PEPTIDES (English)
     Patent Assignee: UNIV TÜLANE
                                             (US)
     Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE; TAYLOR JOHN E:
       KIM SUN HYUK
     Priority (No,Kind,Date): US 257998 A 19881014; 19881209; US 317941 A 19890302; US 376555 A
                                                          19881014; US 282328
                                                                        19890707; US 397169
             19890821; wo 89us4616 w
                                              19891013
     Applic (No, Kind, Date): HK 98111817 A
                                                          19981106
               C07K; A61K
       Abstract No: * 112(17)158978R; 113(19)172755T; 115(15)150377K; 123(21)286737A; 128(18)213739W; 129(02)016394Z
     CA Abstract No: *
     Derwent WPI Acc No: * C 89-309505; C 90-147822; C 95-169633; C 98-229235; C 98-296827; C 99-189718 Language of Document: English
                                                                           91-087241: C
HUNGARY (HU)
  Patent (No, Kind, Date): HU 8906391 AO 19910729
     PEPTIDES WITH MEDICATIVE EFFECT (English)
     Patent Assignee: ADMINISTRATORS OF THE TULANE
    Author (Inventor): COY DAVID; MOREAU JACQUES-PIERRE; TAYLOR JOHN E;
    Priority (No,Kind,Date): US 257998 A 19881014; 19881209; US 317941 A 19890302; US 376555 A
                                                          19881014; US 282328
                                                                        19890707; US 397169
            19890821
    Applic (No,Kind,Date): HU 9163 A
                                                    19891013
    CA Abstract No: * 112(17)158978R; 113(19)172755T
    Derwent WPI Acc No: * C 89-309505; C 90-147822; C 91-087241
    Language of Document: Hungarian
  Patent (No,Kind,Date): HU 90068
THERAPEUTIC PEPTIDES (English)
                                HU 9006872 AO 19910729
    Patent Assignee: ADMINISTRATORS OF THE TULANE
Author (Inventor): COY DAVID; MOREAU JACQUES-PIERRE
Priority (No,Kind,Date): US 394727 A 19890816
Applic (No,Kind,Date): HU 906872 A 19900816
    Derwent WPI Acc No: *
                                 C 91-087240
    Language of Document: Hungarian
 Patent (No,Kind,Date): HU T59420 A2 19920528
PROCESS FOR PRODUCING PEPTIDES HAVING PHARMACEUTICAL ACTION (English)
    Patent Assignee: UNIV TULANE
    Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE; TAYLOR JOHN E;
       KIM SUN HYUK
    Priority (No, Kind, Date): US 257998 A 19881209; US 317941 A 19890302; US
                                                         19881014; US 282328
                                     19890302; US 376555 A
                                                                      19890707; us 397169
            19890821
    Applic (No, Kind, Date): HU 9163 A
                                                    19891013
    IPC: * C07K-007/02; C07K-007/08; C07K-007/06; C07K-007/10; C07K-007/30 CA Abstract No: * 112(17)158978R; 113(19)172755T; 115(15)150377K Derwent WPI Acc No: * C 89-309505; C 90-147822; C 91-087241
    Language of Document: Hungarian
 Patent (No, Kind, Date): HU T65465 A2 19940628
    PROCESS FOR PRODUCING THE RAPEUTIC PEPTIDES (English)
    Patent Assignee: UNIV TULANE (US)
    Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US)
   Priority (No,Kind,Date): US 394727 A 19890816

Applic (No,Kind,Date): HU 906872 A 19900816

IPC: * C07K-005/02; C07K-005/08; C07K-005/06; C07K-005/10; C07K-007/02; C07K-007/08

CA Abstract No: * 115(15)151906U
    Derwent WPI Acc No: * C 91-087240
    Language of Document: Hungarian
 Patent (No, Kind, Date): HU 208439 B
                                                    19931028
    PROCESS FOR PRODUCING PHARMACEUTICAL PEPTIDES (English)
    Patent Assignee: UNIV TULANE
   Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE; TAYLOR JOHN E;
      KIM SUN HYUK
    Priority (No, Kind, Date): US 257998 A
                                                         19881014; US 282328
      19881209; US 317941 A 19890302; US 376555 A
                                                                       19890707; US 397169
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19890821
      Applic (No,Kind,Date): HU 9163 A
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     IPC: * C07K-007/02; C07K-007/08; C07K-007/06; C07K-007/10; C07K-007/30
CA Abstract No: * 112(17)158978R; 113(19)172755T; 115(15)150377K
     Derwent WPI Acc No: * C 89-309505; C 90-147822; C 91-087241 Language of Document: Hungarian
IRELAND (IE)
Patent (No, Kind, Date): IE 91902958 A1 19910227
     SUBSTANCE P ANTAGONISTS (English)
     Patent Assignee: UNIV TULANE (US)
     Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE
     Priority (No, Kind, Date): US 394727 A 19890816
  Applic (No, Kind, Date): IE 902958 A 19900815
IPC: * C07K-007/02; C07K-007/06
CA Abstract No: * 115(15)151906U; 123(21)286737A
Derwent WPI Acc No: * C 91-087240; C 95-169633
Language of Document: English
Patent (No, Kind, Date): IE 9777033 B 19971119
SUBSTANCE DANTACONISTS (English)
     SUBSTANCE P ANTAGONISTS (English)
     Patent Assignee: UNIV TULANE (US)
     Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE
     Priority (No, Kind, Date): US 394727 A
                                                                 19890816
     Applic (No,Kind,Date): IE 902958 A 19900815
IPC: * C07K-007/02; C07K-007/06
CA Abstract No: * 115(15)151906U; 123(21)286737A
Derwent WPI Acc No: * C 91-087240; C 95-169633
Language of Document: English
JAPAN (JP)
  Patent (No, Kind, Date): JP 3141961 A2 19910617
     GAME SET HAVING TWO-FACED PIECE (English)
     Patent Assignee: SUCHIYUAATO EMU RAMURE
Author (Inventor): SUCHIYUAATO EMU RAMURE
     Priority (No,Kind,Date): US 398172 A Applic (No,Kind,Date): JP 90220174 A
                                                                 19890823
                                                                 19900823
                A63F-001/00
     Language of Document: Japanese
  Patent (No, Kind, Date): JP 2795449 B2 19980910
     Patent Assignee: ADOMINISUTOREETAAZU OBU ZA TSU
     Author (Inventor): KOI DEEBITSUDO ETSUCHI; MOROO JATSUKUUPIEERU
     Priority (No,Kind,Date): US 100571 A 19870
Applic (No,Kind,Date): JP 88509311 A 19880
IPC: * C07K-014/46; A61K-038/22; C07K-014/575
                                                                 19870924
                                                                 19880923
  Language of Document: Japanese
Patent (No,Kind,Date): JP 2919889 B2 19990719
     Patent Assignee: ADOMINISUTOREETAAZU OBU ZA TSU
     Author (Inventor): KOI DEIBITSUDO EICHI; MOROO JATSUKUPIEERU; TEIRAA
        JON II; KIMU SUN HYUKU
     Priority (No, Kind, Date): 19881209; US 317941 A
                                           US 257998 A 19881014;
19890302; US 376555 A
                                                                 19881014; US 282328
                                                                                19890707; US 397169
              19890821
     Applic (No, Kind, Date): JP 89511442 A
                                                                 19891013
                CO7K-014/575; CO7K-007/06; A61K-031/00; A61K-038/00;
        A61K-038/04; A61K-038/22
  Language of Document: Japanese
Patent (No,Kind,Date): JP 2502016 T2 19900705
Priority (No,Kind,Date): WO 88US3286 W 198
19870924
                                                                   19880923; US 100571 A
     Applic (No,Kind,Date): JP 88509311 A 19880923 IPC: * C07K-007/06; A61K-037/24; C07K-001/04; C07K-007/08; C07K-099-00 CA Abstract No: * 111(11)097733N
     Derwent WPI Acc No: * C 89-095447
     Language of Document: Japanese
   Patenť (No,Kind,Date): JP 4502922 T2 19920528
     Priority (No, Kind, Date): WO 90US4633 W
                                                                  19900816; US 394727 A
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     Applic (No, Kind, Date): JP 90511667 A
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IPC: * C07K-007/06; A61K-037/02; C07K-099-00 CA Abstract No: * 115(15)151906U
      Derwent WPI Acc No: * C 91-087240
   Language of Document: Japanese Patent (No, Kind, Date): JP 4504406
      Atent (No,Kind,Date): JP 4504406 T2 19920806
Priority (No,Kind,Date): WO 89US4616 W 1989
19881014; US 282328 A 19881209; US 317941
A 19890707; US 397169 A 19890821
Applic (No,Kind,Date): JP 89511442 A 198910
                                                              19891013; US 257998
                                        19881209; US 317941 A
                                                                         19890302; us 376555
                                                           19891013
      IPC: * C07K-007/06; A61K-037/02; A61K-037/24; A61K-037/43; C07K-099-00 CA Abstract No: * 112(17)158978R; 113(19)172755T; 115(15)150377K Derwent WPI Acc No: * C 89-309505; C 90-147822; C 91-087241 Language of Document: Japanese
   Patent (No, Kind, Date): JP 4506664 T2 19921119
      Priority (No, Kind, Date): wo 90us4646 w
                                                             19900817; US 397169
      19890821; US 502438 A 19900330
Applic (No,Kind,Date): JP 90512265
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               C07K-007/06; C05B
      CA Abstract No: * 113(19)172755T; 115(15)150377K
      Derwent WPI Acc No: * C 90-147822; C 91-087241
      Language of Document: Japanese
MONACO (MC)
   Patent (No, Kind, Date): MC 2144
                                                   19920219
      PEPTIDES THERAPEUTIQUES (French)
      Patent Assignee: UNIV TULANE
     Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE; TAYLOR JOHN E;
        KIM SUN HYUK
     Priority (No, Kind, Date): US 257998
                                                          19881014; US 282328
        19881209; wo 89us4616 w
                                           19891013; US 317941 A
                                                                            19890302; US
     376555 A 19890707; US 397169 A Applic (No, Kind, Date): MC 2144 A
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     IPC: *
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     CA Abstract No: * 112(17)158978R; 113(19)172755T; 115(15)150377K;
        123(21)286737A; 128(18)213739W; 129(02)016394z
     Derwent WPI Acc No: * C 89-309505; C 90-147822; C 95-169633; C 98-229235; C 98-296827; C 99-189718 Language of Document: French
                                                                           91-087241; c
  Patent (No, Kind, Date): MC 2193
                                                   19921005
                                             Α
     ENSEMBLE DE JEU COMPORTANT DES PIECES DE JEU A DEUX FACES (French)
     Patent Assignee: STEWART MILTON LAMLE
     Author (Inventor): STEWART MILTON LAMLE
     Priority (No, Kind, Date): US 398172 A
                                                          19890823
     Applic (No,Kind,Date): MC 2144 A
                                                     19900820
               A63F
     Derwent WPI Acc No: * G 91-059703
Language of Document: French
NORWAY (NO)
  Patent (No, Kind, Date): NO 8902060 A
                                                      19890721
     TERAPEUTISKE PEPTIDER. (Norwegian)
     Patent Assignee: UNIV TULANE
                                            (US)
     Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE
     Priority (No,Kind,Date): US 100571 A 19880923
                                                          19870924; wo 88us3286 w
     Applic (No,Kind,Date): NO 892060 A
                                                       19890523
              C07K-007/02; C07K-007/00
     CA Abstract No: *
                             111(11)097733N; 123(21)286737A; 128(18)213739W;
       129(02)016394z
  Derwent WPI Acc No: * C 89-095447; C 95-169633; C 98-229235; C 98-296827; C 99-189718
Language of Document: Norwegian
Patent (No,Kind,Date): NO 9003697 A 19910225
     SPILLSETT MED TOSIDEDE SPILLBRIKKER. (Norwegian)
    Patent Assignee: LAMLE STEWART M
    Author (Inventor): LAMLE STEWART MILTON
    Priority (No, Kind, Date): US 398172 A
    Applic (No, Kind, Date): NO 903697 A
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IPC: * A63F-009/00
  Derwent WPI Acc No: *
                               G 91-059703
  Language of Document: Norwegian
                                                   19920406
Patent (No, Kind, Date): NO 9200678
  TERAPEUTISKE PEPTIDER (Norwegian)
  Patent Assignee: BIOMEASURE INC
                                             (US); UNIV TULANE (US)
  Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE; KIM SUN HYUK Priority (No, Kind, Date): US 397169 A 19890821; US 502438 A 19900330; WO 90US4646 W 19900817 Applic (No, Kind, Date): NO 92678 A 19920220 IPC: * CO7K-005/02; CO7K-007/02; CO7K-005/06; CO7K-005/08; CO7K-005/10; CO7K-007/06; CO7K-007/08; CO7K-007/10
CA Abstract No: * 113(19)172755T; 115(15)150377K; 128(18)213739W Derwent WPI ACC No: * C 90-147822: C 91-087241: C 98-229235: C
  Derwent WPI Acc No: * C 90-147822; C 91-087241; C 98-229235; C
     99-189718
  Language of Document: Norwegian
Patent (No, Kind, Date): NO 8902060
                                              A0 19890523
  TERAPEUTÍSKE PEPTIDER. (Norwegian)
Patent Assignee: UNIV TULANE (US)
  Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE
  Priority (No, Kind, Date): US 100571 A
                                                     19870924; WO 88US3286 W
     19880923
  Applic (No,Kind,Date): NO 892060 A IPC: * CO7K
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  Derwent WPI Acc No: * C 89-095447
Language of Document: Norwegian Patent (No,Kind,Date): NO 9003697
                                              A0 19900822
  SPILLSETT MED TOSIDEDE SPILLBRIKKER. (Norwegian)
  Patent Assignee: LAMLE STEWART MILTON
  Author (Inventor): LAMLE STEWART MILTON
  Priority (No, Kind, Date): US 398172 A
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  Applic (No,Kind,Date): NO 903697 A IPC: * A63F
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Language of Document: Norwegian Patent (No,Kind,Date): NO 9200678
                                              A0 19920220
  TERAPEUTISKE PEPTIDER (Norwegian)
  Patent Assignee: BIOMEASURE INC (US); UNIV TULANE (US)
  Author (Inventor): COY DAVID H; MOREAU JACQUES-PIERRE; KIM SUN HYUK
  Priority (No,Kind,Date): US 397169 A 19900330; WO 90US4646 W 19900817
                                                       19890821; US 502438 A
  Applic (No, Kind, Date): NO 92678 A IPC: * C07K-005/02
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  IPC: * C07K-005/02
CA Abstract No: * 113(19)172755T; 115(15)150377K
Derwent WPI Acc No: * C 90-147822; C 91-087241
Language of Document: Norwegian
Patent (No, Kind, Date): NO 178306 B
                                                  19951120
  ANALOGIFREMGANGSMAATE VED FREMSTILLING AV BOMBESIN-ANTAGONISTPEPTID
     (Norwegian)
  Patent Assignee: UNIV TULANE (US)
  Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE
  Priority (No, Kind, Date): US 100571 A
                                                       19870924; wo 88us3286 w
     19880923
  Applic (No, Kind, Date): NO 892060 A
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  IPC: * C07K-007/06; C07K-007/08
CA Abstract No: * 111(11)097733N; 123(21)286737A; 128(18)213739W;
     129(02)016394Z
  Derwent WPI Acc No: * C 89-095447; C 95-169633; C 98-229235; C
     98-296827; c 99-189718
  Language of Document: Norwegian
Patent (No, Kind, Date): NO 302619 B1 19980330
  ANALOGIFREMGANGSMAATE FOR FREMSTILLING AV ET TERAPEUTISK PEPTID
     (Norwegian)
  Patent Assignee: BIOMEASURE INC (US); UNIV TULANE (US)
  Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE
                                                                                  (US): KIM
     SUN HYUK (US)
  Priority (No, Kind, Date): US 397169 A
                                                       19890821; US 502438 A
     19900330; wo 90us4646 w
                                       19900817
  Applic (No,Kind,Date): NO 92678 A
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IPC: * C07K-005/02
         CA Abstract No: * 113(19)172755T; 115(15)150377K; 128(18)213739W
          Derwent WPI Acc No: * C 90-147822; C 91-087241; C 98-229235; C
              99-189718
          Language of Document: Norwegian
     Patent (No, Kind, Date): NO 178306 C
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         ANALOGIFREMGANGSMAATE VED FREMSTILLING AV BOMBESIN-ANTAGONISTPEPTID
              (Norwegian)
         Patent Assignee: UNIV TULANE (US)
         Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US)
         Priority (No, Kind, Date): US 100571 A 19880923
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         Applic (No,Kind,Date): NO 892060 A IPC: * C07K-007/06; C07K-007/08
                                                                                              19890523
                        C07K-007/06; C07K-007/08
         CA Abstract No: * 111(11)097733N; 123(21)286737A
         Derwent WPI Acc No: * C 89-095447; C 95-169633
Language of Document: Norwegian
NEW ZEALAND (NZ)
     Patent (No, Kind, Date): NZ 234993 A
                                                                                         19920428
         STACKABLÉ INDICIA BEARING PIECES AS GAME SET (English)
         Patent Assignee: LAMLE STEWART M
         Author (Inventor): LAMLE STEWART MILTON
        Priority (No,Kind,Date): US 398172 A
Applic (No,Kind,Date): NZ 234993 A
IPC: * A63F-001/02; A63F-009/20
Derwent WPI Acc No: * G 91-059703
                                                                                                  19890823
                                                                                             19900821
        Language of Document: English
PORTUGAL (PT)
    Patent (No, Kind, Date): PT 95016 A
                                                                                      19910418
        PROCESSO PARA A PREPARACAO DE PEPTIDOS TERAPEUTICOS (English; French;
            German; Portugese)
        Patent Assignee: UNIV TULANE
        Priority (No, Kind, Date): US 394727
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        Applic (No, Kind, Date): PT 95016 A
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        IPC: *
                         C07K-001/00
   Derwent WPI Acc No: * C 91-087240
Language of Document: Portugese
Patent (No, Kind, Date): PT 95057 A
                                                                                      19910522
        PROCESSO PARA A PREPARACAO DE PEPTIDOS TERAPEUTICOS (English; French;
            German; Portugese)
       Patent Assignee: BIOMEASURE INC (US); UNIV TULANE (US)
       Author (Inventor):
                                                  COY DAVID HOWARD (US); MOREAU JACQUES-PIERRE (US)
       ; KIM SUN HYUK (US)
Priority (No, Kind, Date): US 397169 A
Applic (No, Kind, Date): PT 95057 A

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       IPC: * C07K-007/08; A61K-037/02
CA Abstract No: * 113(19)172755T
       Derwent WPI Acc No: * C 90-147822; C 91-087241
       Language of Document: Portugese
   Patent (No, Kind, Date): PT 95016 B
                                                                                     19970528
       PROCESSO PARA A PREPARACAO DE PEPTIDOS TERAPEUTICOS (English; French;
            German; Portugese)
       Patent Assignee: UNIV TULANE (US)
Priority (No, Kind, Date): US 394727 A 19890816
Applic (No, Kind, Date): PT 95016 A 19900816
IPC: * CO7K-007/02; CO7K-007/22; A61K-038/08
CA Abstract No: * 115(15)151906U; 123(21)286737A
                                                                                                19890816
       Derwent WPI Acc No: * C 91-087240; C 95-169633
Language of Document: Portugese
   Patent (No,Kind,Date): PT 95057 B
                                                                                     19971231
       PROCESSO PARA A PREPARACAO DE PEPTIDOS TERAPEUTICOS (English; French;
           German; Portugese)
       Patent Assignee: BIOMEASURE INC (US); UNIV TULANE (US)
Author (Inventor): COY DAVID HOWARD (US); MOREAU JACQUES-PIERRE (US)
               KIM SUN HYUK (US)
       Priority (No, Kind, Date): US 397169 A
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Applic (No,Kind,Date): PT 95057 A
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                 C07K-007/02; C07K-007/06; C07K-007/08; C07K-014/595
      CA Abstract No: * 113(19)172755T; 115(15)150377K
      Derwent WPI Acc No: * C 90-147822; C 91-087241
      Language of Document: Portugese
RUSSIA (RU)
   Patent (No,Kind,Date): RU 2088592 C1 19970827
      THERAPEUTIC PEPTIDES OR THEIR PHARMACEUTICALLY ACCEPTABLE SALTS
         (English)
      Patent Assignee: UNIV TULANE (US)
      Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US);
         TAYLOR JOHN E (US); KIM SUN HYUK
                                                           (US)
      Priority (No, Kind, Date): WO 89US4616 W
                                                                     19891013; US 317941 A
      19890302; US 376555 A 19890707; US 397169 A Applic (No, Kind, Date): RU 4895537 A 19891013
                                                                                  19890821
                 C07K-007/06; A61K-038/08
     CA Abstract No: * 113(19)172755T; 115(15)150377K; 123(21)286737A Derwent WPI Acc No: * C 90-147822; C 91-087241; C 95-169633 Language of Document: Russian
SLOVAKIA (SK)
   Patent (No, Kind, Date): SK 9004028 A3 20000711
      SUBSTANCE P ANALOG (English)
      Patent Assignee: UNIV TULANE (US)
     Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US) Priority (No, Kind, Date): US 394727 A 19890816 Applic (No, Kind, Date): SK 904028 A 19900816 IPC: * CO7K-007/02; CO7K-007/22; A61K-038/08 CA Abstract No: * 115(15)151906U; 123(21)286737A
     Derwent WPI Acc No: * C 91-087240; C 95-169633
      Language of Document: Slovak
  Patent (No, Kind, Date): SK 280796 B6 20000711
      SUBSTANCE P ANALOG (English)
     Patent Assignee: UNIV TULANE (US)
Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US)
Priority (No,Kind,Date): US 394727 A 19890816
Applic (No,Kind,Date): SK 904028 A 19900816
     IPC: * C07K-007/02; C07K-007/22; A61K-038/08
CA Abstract No: * 115(15)151906U; 123(21)286737A
Derwent WPI Acc No: * C 91-087240; C 95-169633
Language of Document: Slovak
UNITED STATES OF AMERICA (US)
Patent (No, Kind, Date): US 4998737 A
                                                             19910312
      TWO-SIDED PLAYING PIECE GAME SET (English)
      Patent Assignee: LAMLE STEWART M (US)
     Author (Inventor): LAMLE STEWART M (US) Priority (No, Kind, Date): US 398172 A
                                                                 19890823
     Applic (No, Kind, Date): US 398172 A National Class: * 273296000; 273292000
                                                               19890823
  IPC: * A63F-001/00
Language of Document: English
Patent (No,Kind,Date): US 5084555 A
                                                             19920128
     AN OCTAPEPTIDE BOMBESIN ANALOG (English)
Patent Assignee: UNIV TULANE (US); BIOMEASURE INC (US)
      Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US); KIM
        SUN H (US)
     Priority (No,Kind,Date): US 397169 B2 19890821; US 376555 B2 19890707; US 317941 B2 19890302; US 282328 A2 19881209; US 257998 B2 19881014; US 248771 B2 19880923; US 207759 B2 19880616; US 204171 B2 19880608; US 173311 B2 19880325; US 100571 B2
        19870924
      Applic (No, Kind, Date): US 502438 A
                                                               19900330
      National Class: * 530328000; 530309000; 530323000; 530324000;
         530325000; 530326000; 530327000; 530329000; 530332000
      IPC: * C07K-007/06; C07K-007/30
      CA Abstract No: * 111(11)097733N; 112(17)158978R; 112(19)179890W;
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113(19)172755T; 115(15)150377K
    Derwent WPI Acc No: * C 89-095447; C 89-280003; C 89-309505; C
       90-147822; c 91-087241
    Language of Document: English
 Patent (No, Kind, Date): US 5162497 A
                                                           19921110
    BRADYKININ ANALOGS WITH NON-PEPTIDE BOND (English)
    Patent Assignee: UNIV TULANE (US)
    Author (Inventor); COY DAVID H (US); MOREAU JACQUES-PIERRE (US);
    TAYLOR JOHN E (US); KIM SUN H (US)
Priority (No, Kind, Date): US 257998 B2 19881014; US 248771 B2
19880923; US 207759 B2 19880616; US 204171 B2 19880608; US 173311
       B2 19880325; US 100571 B2 19870924
    Applic (No, Kind, Date): US 282328 A
                                                            19881209
    National Class: * 530314000; 530332000; 530328000; 514803000;
       930030000; 930DIG790; 930DIG600; 930DIG601
C: * C07K-007/00; C07K-007/18
                              111(11)097733n; 112(17)158978R; 112(19)179890w;
    CA Abstract No: *
       113(19)172755T
    Derwent WPI Acc No: * C 89-095447; C 89-280003; C 89-309505; C
       90-147822
   Language of Document: English
 Patent (No, Kind, Date): US 5410019 A THERAPEUTIC PEPTIDES (English)
                                                          19950425
   Patent Assignee: UNIV TULANE (US)
Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US)
Priority (No, Kind, Date): US 860675 A 19920330; US 394727 B2
19890816; US 317941 B2 19890302; US 282328 A2 19881209; US 2579
B2 19881014; US 248771 B2 19880923; US 207759 B2 19880616; US
204171 B2 19880608; US 173311 B2 19880325; US 100571 B2
                                                                              19881209; US 257998
      19870924
   Applic (No, Kind, Date): US 860675 A
                                                            19920330
   Addnl Info: 5162497 Patented
   National Class: * 530323000; 530327000; 530328000; 530329000;
      530330000
  CA Abstract No: * 111(11)097733N; 112(17)158978R; 112(19)179890W; 113(19)172755T; 115(15)151906U; 123(21)286737A; 128(18)213739W; 129(02)016394Z; 123(21)286737A

Derwent WPI Acc No: * C 89-095447; C 89-280003; C 89-309505; C 90-147822; C 91-087240; C 95-169633; C 98-229235; C 98-22682
              C07K-007/02; C07K-007/06
                        No: * C 89-095447; C 89-280003; C 89-309505; C 91-087240; C 95-169633; C 98-229235; C 98-296827; C
      99-189718; c 95-169633
   Language of Document: English
Patent (No, Kind, Date): US 5723578 A
                                                          19980303
   Peptide analogs of bombesin (English)
   Patent Assignee: ADMINISTRATORS OF TULANE EDUCA (US); BIOMEASURE INC
      (US)
   Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US); KIM
      SUN HYUK (US)
   Priority (No,Kind,Date): US 488099 A 19950607; US 337127 19941110; US 779039 B2 19911018; US 502438 A2 19900330;
                                                                             19900330; us 397169
          19890821; US 376555
                                        B2 19890707; US 317941
                                                                              В2
                                                                                   19890302; US
      282328 A2 19881209; US 257998 B2 19881014; US 248771 19880923; US 207759 B2 19880616; US 204171 B2 1988060
                                                                            19880608; us 173311
  B2 19880325; US 100571 B2 19870924

Applic (No,Kind,Date): US 488099 A 19950607

Addnl Info: 5084555 Patented; 5162497 Patented

National Class: * 530326000; 530327000; 530328000

IPC: * A61K-038/00; C07K-005/00; C07K-007/00; C07K-017/00
  Derwent WPI Acc No: ; C 98-229235
  Language of Document: English
Patent (No, Kind, Date): US 5750646 A
                                                         19980512
   BRADYKININ ANALOGS WITH NON-PEPTIDE BOND (English)
  Patent Assignee: UNIV TULANE (US); BIOMEASURE INC (US)
  Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US);
     TAYLOR JOHN E (US); KIM SUN HYUK (US)
  Priority (No, Kind, Date): US 408197 A 19950322; US 880179 B2 19920507; US 282328 A2 19881209; US 257998 B2 19881014; US 248771
     B2 19880923; US 207759 B2 19880616; US 204171 B2 19880608; US
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173311 B2 19880325; US 100571 A2
                                                                      19870924
   Applic (No, Kind, Date): US 408197
                                                                      19950322
                                                               Α
   Addnl Info: 5162497 Patented National Class: * 530314000; 530328000; 530332000
   IPC: * A61K-038/00; C07K-005/00; C07K-007/00; C07K-017/00
   CA Abstract No: * 111(11)097733N; 112(17)158978R; 112(19)179890W;
       113(19)172755T; 123(21)286737A; 128(18)213739W; 129(02)016394Z;
       129(02)016394Z
   Derwent WPI Acc No: * C 89-095447; C 89-280003; C 89-309505; C 90-147822; C 95-169633; C 98-229235; C 98-296827; C 99-189718; C
       98-296827
   Language of Document: English
Patent (No, Kind, Date): US 5877277
   OCTAPEPTIDE BOMBESIN ANALOGS (English)
   Patent Assignee: BIOMEASURE INC (US); UNIV TULANE (US)
   Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US); KIM
       SUN HYUK (US)
   Priority (No,Kind,Date): US 337127 A 19941110; US 779039 B1 19911018; US 502438 A2 19900330; US 397169 B2 19890821; US 376555 B2 19890707; US 317941 B2 19890302; US 282328 A2 19881209; US 257998 B2 19881014; US 248771 B2 19880923; US 207759 B2 19880616; US 204171 B2 19880608; US 173311 B2 19880325; US 100571
            19870924
   Applic (No, Kind, Date): US 337127 A
                                                                     19941110
  Addnl Info: 5084555 Patented; 5162497 Patented
National Class: * 530328000; 530323000
IPC: * C07K-005/00; C07K-007/06; A61K-038/00
CA Abstract No: * 111(11)097733N; 112(17)158978R; 112(19)179890W; 113(19)172755T; 115(15)150377K; 123(21)286737A; 128(18)213739W;
       129(02)016394Z
   Derwent WPI Acc No: * C 89-095447; C 89-280003; C 89-309505; C
       90-147822; C
                              91-087241; C 95-169633; C 96-455920; C 98-229235; C
                                                    99-189718
       98-296827; C 99-189718; C
   Language of Document: English
                                                                   AA 20030313
Patent (No, Kind, Date): US 20030050436
   Octapeptide bombesin analogs (English)
Patent Assignee: BIOMEASURE INC MASSACHUSETTS C (US)
   Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US); KIM
      SUN HYUK (US)
   Priority (No, Kind, Date): US 4530 A
                                                                    20011023; US 260846 A3
      19990302; ús 337127 A3 19941110; us 779039 B1 19911018; us 502438
  19990302; US 337127 A3 19941110; US 779039 B1 19911018; US 3024.
A2 19900330; US 397169 B2 19890821; US 376555 B2 19890707; US
317941 B2 19890302; US 282328 A2 19881209; US 257998 B2
19881014; US 248771 B2 19880923; US 207759 B2 19880616; US 2041
B2 19880608; US 173311 B2 19880325; US 100571 B2 19870924
Applic (No,Kind,Date): US 4530 A 20011023
Addnl Info: 6307017 Patented; 5877277 Patented; 5084555 Patented;
                                                                                           19880616; US 204171
       5162497 Patented
   National Class: * 530328000: 530329000
   IPC: * C07K-007/08; C07K-007/06
Derwent WPI Acc No: ; C 03-810756
Language of Document: English
Patent (No,Kind,Date): US 6307017
                                                                   20011023
                                                             BA
   Octapeptide bombesin analogs (English)
Patent Assignee: BIOMEASURE INC (US); UNIV TULANE (US)
   Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE
                                                                                                             (US); KIM
      SUN HYUK (US)
  Priority (No,Kind,Date): US 260846 A 19990302; US 337127 A3 19941110; US 779039 B2 19911018; US 502438 A2 19900330; US 397169 B2 19890821; US 376555 B2 19890707; US 317941 B2 19890302; US 282328 A2 19881209; US 257998 B2 19881014; US 248771 B2 19880923; US 207759 B2 19880616; US 204171 B2 19880608; US 173311 B2 19880325; US 100571 B2 19870924 Applic (No,Kind,Date): US 260846 A 19990302
   Applic (No, Kind, Date): US 260846 A 19990302
Addnl Info: 5877277 Patented; 5084555 Patented; 5162497 Patented
National Class: * 530328000; 530300000; 530323000; 514012000;
       514015000
   IPC: * A61K-038/00; A61K-038/04; C07K-005/00; C07K-007/00
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Derwent WPI Acc No: ; C 02-162970 Language of Document: English WORLD INTELLECTUAL PROPERTY ORGANIZATION, PCT (WO) Patent (No, Kind, Date): WO_8902897 A1 19890406 THERAPEUTIC PEPTIDES (English) Patent Assignee: UNIV TULANE (US); COY DAVID HOWARD (US); MOREAU Author (Inventor): CO Priority (No " Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US) Priority (No, Kind, Date): US 100571 A 19870924 Applic (No, Kind, Date): WO 88US3286 A 19880923 Designated States: (National) AU; DK; FI; JP; NO (Regional) AT; BE; CH; DE; FR; GB; IT; LU; NL; SE Filing Details: WO 13000 With international search report; Before expiration of time limit for amending the claims and to be republished in the event of the receipt of the amendments IPC: * C07K-007/02; C07K-007/06; C07K-007/08 Language of Document: English Patent (No, Kind, Date): WO 8909230 A1 19891005 THERAPEUTIC PEPTIDES (English) Patent Assignee: UNIV TULANE (US) Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US); TAYLOR JOHN E (US)
Priority (No, Kind, Date): US 173311 A 19880325
Applic (No, Kind, Date): WO 89US1216 A 19890322
Designated States: (National) AU; DK; FI; JP; NO
Filing Details: WO 13000 With international search report; Before expiration of time limit for amending the claims and to be republished in the event of the receipt of the amendments C07K - 007/18Language of Document: English Patent (No, Kind, Date): WO 8909231 A1 19891005 THERAPEUTIC PEPTIDES (English) Patent Assignee: UNIV TÜLANE (US) Author (Inventor): COY DAVID H (US); MO TAYLOR JOHN E (US); KIM SUN HYUK (US) (US); MOREAU JACQUES-PIERRE Priority (No, Kind, Date): US 173311 A 19880325; US 282328 19881209 Applic (No,Kind,Date): WO 89US1259 A 19890327 Designated States: (National) AU; DK; FI; JP; NO (Region CH; DE; FR; GB; IT; LU; NL; SE Filing Details: WO 10000 With international search report (Regional) AT; BE; C07K-007/18 CA Abstract No: ; 112(17)158978R Derwent WPI Acc No: ; C 89-309505 Language of Document: English Patent (No, Kind, Date): WO 9003980 A1 19900419 THERAPEUTIC PEPTIDES (English)
Patent Assignee: UNIV TULANE (US) Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (us) TAYLOR JOHN E (US); KIM SUN HYUK Priority (No, Kind, Date): US 257998 A 19881014; US 282328 19881209; ÚS 317941 A 19890302; US 376555 A 19890707; us 397169 19890821 Applic (No, Kind, Date): WO 89US4616 A 19891013 Designated States: (National) AU; BB; BG; BR; DK; FI; HU; JP; KP; KR; LK; MC; MG; MW; NO; RO; SD; SU (Regional) AT; BE; BF; BJ CH; CM; DE; FR; GA; GB; IT; LU; ML; MR; NL; SE; SN; TD; TG Filing Details: WO 10000 With international search report (Regional) AT; BE; BF; BJ; CF; CG; C07K-007/02; C07K-007/06; C07K-007/08; C07K-007/10; C07K-007/30 CA Abstract No: ; 113(19)1727557 Derwent WPI Acc No: ; C 90-147822 Language of Document: English Patent (No, Kind, Date): WO 9102745 A1 19910307 THERAPEUTIC PEPTIDES (English) Author (Inventor): COY DAVID H (UPriority (No.Kind Date): 12 2011 Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US) Priority (No, Kind, Date): US 394727 A 19890816

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Applic (No,Kind,Date): WO 90US4633 A
                                                            19900816
   Designated States: (National) CA; HU; JP
                                                                (Regional) AT; BE; CH; DE;
   DK; ES; FR; GB; IT; LU; NL; SE
Filing Details: WO 130000 With international search report; Before
     expiration of time limit for amending the claims and to be
     republished in the event of the receipt of the amendments
  IPC: * C07K-005/02; C07K-005/06; C07K-005/08; C07K-005/10; C07K-007/02
; C07K-007/06; C07K-007/08
  CA Abstract No: ; 115(15)151906U
Derwent WPI Acc No: ; C 91-087240
Language of Document: English
Patent (No,Kind,Date): WO 9102746 A1 19910307
THERAPEUTIC PEPTIDES (English)
  Patent Assignee: BIOMEASURE INC (US); UNIV TULANE (US)
  Author (Inventor): COY DAVID H (US); MOREAU JACQUES-PIERRE (US); KIM
     SUN HYUK (US)
   Priority (No, Kind, Date): US 397169 A
                                                            19890821; US 502438 A
     19900330
  Applic (No, Kind, Date): WO 90US4646 A
                                                            19900817
  Designated States: (National) AU; CA; FI; JP; NO (Regional) AT; B CH; DE; DK; ES; FR; GB; IT; LU; NL; SE Filing Details: WO 130000 With international search report; Before
                                                                            (Regional) AT; BE;
  Filing Details: WO 130000 With international search report expiration of time limit for amending the claims and to be
     republished in the event of the receipt of the amendments
  IPC: * C07K-005/02; C07K-005/06; C07K-005/08; C07K-005/10; C07K-007/02; C07K-007/06; C07K-007/08; C07K-007/10; C07K-007/30

CA Abstract No: ; 115(15)150377K

Derwent WPI Acc No: ; C 91-087241
  Language of Document: English
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9/17/2004

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the non-peptide bond, failing to exhibit the <u>in vivo</u> activity of naturally occurring bombesin. (A detailed discussion of the chemistry of non-peptide bonds is given in Coy et al. (1988) Tetrahedron <u>44</u>,3:835-841, hereby incorporated by reference.)

Preferably, naturally occurring bombesin is 5 characterized in that one or more amino acids in the amino terminal half of bombesin are hydrogen bonded to one or more amino acids in the carboxy terminal half of bombesin, and the non-peptide bond of the linear peptide decreases that hydrogen bonding, thereby destroying biological activity. It is 10 believed that many of the linear peptides of the invention are analogs of bombesin whose biological activity depends at least in part on their ability to form tertiary "hairpin" configurations in which amino acids in the amino terminal ("left") half of the molecule are hydrogen bonded to amino 15 acids in the carboxy terminal ("right") half of the molecule, and that the pseudopeptide bond introduced according to the invention interferes with this hydrogen bonding, hindering the formation of the hairpin configuration on which activity depends. One may expect the loss of the ability to hydrogen 20 bond to affect the biological activity of the molecule either by the loss of structural stability conferred by the transannular bonding or by the inability of the backbone to hydrogen bond to the receptor. Additionally, the increased flexibility of the molecule about the reduced bond compared 25 with the rigidity of the normal peptide amide bond is expected to alter the conformational integrity of the molecule and thus its biological activity.

It is apparent from the above that the linear peptides
for which introduction of a pseudopeptide bond is useful in
creating or enhancing antagonist activity are those in which
activity is associated with a site within the amino acid chain
(some peptides, e.g., CCK, have their active sites at an end of
the peptide). The pseudopeptide bond can be introduced in a

region involved in receptor binding, or in a non-binding region; it has been shown (Nagain et al., Peptides, 8:1023-28 (1987)) that a pseudopeptide bond introduced in the binding region does not prevent binding. Generally, useful classes of peptides in which this modification can be made are those in which at least one amino acid involved in the active site is located in the carboxy terminal half of the molecule; the non-peptide bond is introduced between this amino acid and one adjacent to it.

One class of peptides of the invention is an effective bombesin antagonist peptide of formula (1):

wherein

15 $A^1 = pGlu \text{ or is deleted};$

A² = Gln, Asn, Gly, Ala, Leu, Ile, Nle, α-aminobutyric acid, Met, Val, Phe, p-X-Phe (X = F, Cl, Br, OH or CH₃), Trp, β-naphthylalanine or is deleted;

20 A_{4}^{3} = Arg, D-Arg, Lys, D-Lys or is deleted;

A⁴ = Gln, Asn, Gly, Ala, Leu, Ile, Nle,
α-aminobutyric acid, Met, Val, Phe, p-X-Phe
(X = F, Cl, Br, OH or CH₃), Trp,
β-naphthylalanine or is deleted;

25 A⁵ = Gln, Asn, Gly, Ala, Leu, Ile, Nle, α-aminobutyric acid, Met, Val, Phe, D-Phe, p-X-Phe (X = F, Cl, Br, OH or CH₃), Trp, β-naphthylalanine, D-Ala or is deleted;

- A^6 = Gln, Asn, Gly, Ala, D-Ala, N-Ac-D-Ala, Leu, Ile, Nle, α -aminobutyric acid, Met, Val, Phe, p-X-Phe(X = F, Cl, Br, OH or CH₃), Trp, p-Glu, ß-naphthylalanine or is deleted;
- 5 A^7 = Gln, Asn, Gly, Ala, Leu, Ile, Nle, α -aminobutyric acid, Met, Val, Phe, D-Phe, p-X-Phe (X = F, Cl, Br, OH or CH₃), Trp, His, or β -naphthylalanine;
 - $A^8 = Trp;$
- 10 A^9 = Gln, Asn, Gly, Ala, Leu, Ile, Nle, α -aminobutyric acid, Met, Val, Phe, p-X-Phe (X = F, Cl, Br, OH or CH₃), Trp, or β -naphthylalanine;
- $A^{10} = Gln$, Asn, Gly, Ala, Leu, Ile, Nle, α -aminobutyric acid, Met, Val, Phe, p-X-Phe (X = F, Cl, Br, OH or CH₃), Trp, or β -naphthylalanine;
 - $A^{11} = Gly, or D-Ala;$
 - A^{12} = His, Phe, or p-X-Phe (X = F, C1, Br, OH, CH₃);
- 20 $A^{13} = Gln$, Asn, Gly, Ala, Leu, Ile, Nle, α -aminobutyric acid, Met, Val, Phe, p-X-Phe (X = F, Cl, Br, OH or CH₃), Trp, β -naphthylalanine;
- $A^{14} = Gln$, Asn, Gly, Ala, Leu, Ile, Nle, α -aminobutyric acid, Met, Val, Phe, p-X-Phe (X = F, Cl, Br, OH or CH₃), Trp, or β -naphthylalanine;

provided that

each R_1 , R_2 , R_3 and R_4 , independently, is H, C_{1-12} alkyl, C_{7-10} phenylalkyl, COE_1 (where E_1 is C_{1-20} alkyl, C_{3-20} alkenyl, C_{3-20} alkinyl, phenyl, naphthyl, or C_{7-10} phenylalkyl), or $COOE_2$ (where E_2 is C_{1-10} alkyl or C_{7-10} phenylalkyl), and R_1 and R_2 are bonded to the N-terminal amino

acid of said peptide, which can be A^1 , A^2 , A^3 , A^4 , A^5 , A^6 , or A^7 , provided that when one of R₁ or R₂ is COE₁ or COOE₂, the other must be H, and when one of R_3 or R_4 is COE_1 or $COOE_2$, the other must be H, and further provided that when A^1 = pGlu, \mathbf{R}_1 must be H and \mathbf{R}_2 must be the portion of Glu that forms the imine ring in pGlu; and for each of the residues A^7 , A^8 , A^9 , A^{10} , A^{11} , A^{12} , and A¹³, independently, the carbon atom participating in the amide bond between that residue and the nitrogen atom of the alpha amino group of the adjacent amino acid 10 residue may be a carbonyl carbon or may be reduced to a methylene carbon, provided that at least one such carbon atom must be reduced to a methylene carbon (i.e., at least one of the subject peptide CONH bonds must be replaced by a non-peptide, i.e., pseudopeptide, CH2NH 15 bond); or a pharmaceutically acceptable salt thereof. (Where no D- or L-isomeric designation is given herein, the naturally occurring L-isomer is intended.)

Preferably, an effective bombesin antagonist peptide has, for each of the residues A¹¹, A¹², and 20 ${\tt A}^{13}$, independently, the carbon atom participating in the amide bond between that residue and the nitrogen atom of the alpha amino group of the adjacent amino acid residue which may be a carbonyl carbon or may be reduced to a methylene carbon, provided that at least one such 25 carbon atom must be reduced to a methylene carbon; or a pharmaceutically acceptable salt thereof. Most preferably, the bombesin antagonist peptide has ${\tt A}^1$ through A^6 deleted and the carbon atom participating in the amide bond between Leu^{13} and Leu^{14} is a 30 methylene carbon, or a pharmaceutically acceptable salt thereof

Another class of peptides of the invention are bombesin-related antagonist peptides derived from litorin and of the amino acid formula:

R₁ A¹ -A² -A³ -A⁴ -A⁵ -A⁶ -A⁷ -A⁸ -A⁹ R₃

wherein A¹ is pGlu; A² is Gln; A³ is Trp; A⁴ is Ala; A⁵ is Val; A⁶ is Gly or D-Ala; A⁷ is His; A⁸ is Phe or Leu; and A⁹ is Met or Leu; provided that the carbon atom participating in the amide bond between the A⁸ residue and the nitrogen atom of the alpha amino group of the adjacent amino acid residue may be a carbonyl carbon or may be reduced to a methylene carbon, or a pharmaceutically acceptable salt thereof.

Peptides of the invention that contain a pseudopeptide bond substitution within the active site of the naturally occurring peptide are antagonists to 15 the biological activity of the naturally occurring bombesin peptide, with one exception which we have observed; the linear analog of bombesin BIM-26027 $[Val^{10}\psi[CH_2NH]Leu^{14}]BN$ is an agonist of the biological activity of naturally occurring bombesin. 20 (Non-peptide bonds are symbolized herein by " ψ [CH₂NH]" or " ψ ".) Therefore, a third class of peptides of the invention are effective bombesin agonists of the formula (1) recited above, including, for each of the residues A^9 , A^{10} , A^{11} , A^{12} , 25 A^{13} , and A^{14} , independently, the carbon atom participating in the amide bond between that residue and the nitrogen atom of the alpha amino group of the adjacent amino acid residue may be a carbonyl carbon or may be a non-peptide bond, provided that the non-peptide 30 bond may be a carbonyl carbon having been reduced to a methylene carbon; further provided that at least one

such carbon atom must be reduced to a methylene carbon; or a pharmaceutically acceptable salt thereof. Most preferred is the bombesin agonist having the formula pGlu-Gln-Arg-Leu-Gly-Asn-Gln-Trp-Ala-Val-Gly-His-Leu-Leu[Val¹⁰Y[CH₂NH]Leu¹⁴].

Other agonist analogues are peptides in which either the pseudopeptide bond is not located in the active site of the naturally occurring peptide, or in which two amino acid residues of the active site are replaced by statine or AHPPA.

(Statine has the chemical structure

and statine-amide has the structure

15 and AHPPA has the formula:

(3S,4S)-4-amino-3-hydroxy-5-phenylpentanoic acid.)
Therefore, a fourth class of peptides of the invention is an effective bombesin agonist which is an analog of naturally occurring, biologically active bombesin having an active site, which includes positions A⁹, A¹⁰, A¹¹, A¹², A¹³, and A¹⁴, and a binding site responsible for the binding of bombesin to a receptor on a target cell, the analog having either (a) a non-peptide bond outside of the active site of bombesin, or (b) having at least one statine or AHPPA residue in place of two naturally occurring amino acids of the active site; and further, the peptide can contain statine or AHPPA when all bonds between amino acid

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residues are peptide bonds and, further, when an amino acid residue is statine or AHPPA, the amino acid to the right of it in the formula is deleted, so that the analog is capable of binding to the receptor and, by virtue of the statine or AHPPA residue, exhibiting enhanced in vivo biological activity compared to naturally occurring bombesin. Most preferred in this class is the bombesin agonist having the amino acid formula pGlu-Gln-Arg-Leu-Gly-Asn-Gln-Trp-Ala-Val-Gly-His-[Sta¹³,Des Met¹⁴].

The bombesin antagonists and agonists of the invention are suitable for the treatment of all forms of cancer where bombesin-related substances act as autocrine or paracrine mitotic factors, especially pancreas and small-cell lung carcinoma.

In formula (1), when R_1 , R_2 , R_3 or R_4 is an aromatic, lipophilic group, the <u>in vivo</u> activity can be long lasting, and delivery of the compounds of the invention to the target tissue (e.g., the lungs) can be facilitated.

Other features and advantages of the invention will be apparent from the following description of the preferred embodiments thereof, and from the claims.

Description of the Preferred Embodiments
We will first briefly describe the table.

Table

Table I shows formulas for the pseudo-peptide analogues and results of <u>in vitro</u> inhibition of [125 I]GRP binding to cerebral cortical and 3T3 bombesin receptors, and bombesin-stimulated [3 H]Thymidine uptake by cultured 3T3 cells.

We now describe the structure, synthesis, and use of the preferred embodiments of the invention. Structure

The peptides of the invention all have a non-peptide bond in at least one of the indicated position, except for the statine or AHPPA substituted analogs, such as sta¹³-des Met¹⁴ bombesin. non-peptide bond is meant that the carbon atom participating in the bond between two residues is reduced from a carbonyl carbon to a methylene carbon. The peptide bond reduction method which yields this non-peptide bond is described in Coy et al., U.S. patent application, Serial No. 879,348, assigned to the same 10 assignee as the present application, hereby incorporated by reference. Any one or all of the amino acids in positions 1 through 6 of the bombesin antagonists may be deleted from the peptides, and the peptides are still active as antagonists or agonists.

The peptides of the invention can be provided in the form of pharmaceutically acceptable salts. Examples of preferred salts are those with therapeutically acceptable organic acids, e.g., acetic, lactic, maleic, citric, malic, ascorbic, succinic, 20 benzoic, salicylic, methanesulfonic, toluenesulfonic, or pamoic acid, as well as polymeric acids such as tannic acid or carboxymethyl cellulose, and salts with inorganic acids such as the hydrohalic acids, e.g., 25 hydrochloric acid, sulfuric acid, or phosphoric acid. Synthesis of Bombesin Antagonists

The synthesis of the bombesin antagonist pGlu-Gln-Arg-Leu-Gly-Asn-Gln-Trp-Ala-Val-Gly-His-Leuw[CH2-NH]Leu-NH2 follows. Other bombesin 30 antagonists and agonists can be prepared by making appropriate modifications of the following synthetic method.

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The first step is the preparation of the intermediate pGlu-Gln-Arg(tosyl)-Leu-Gly-Asn-Gln-Trp-Ala-Val-Gly-His(benzyloxycarbonyl)-Leu ψ [CH2NH] Leu-benzhydrylamine resin, as follows.

Benzhydrylamine-polystyrene resin (Vega Biochemicals, Inc.) (0.97 g, 0.5 mmole) in the chloride ion form is placed in the reaction vessel of a Beckman 990B peptide synthesizer programmed to perform the following reaction cycle: (a) methylene chloride; (b) 33% trifluoroacetic acid (TFA) in methylene chloride (2 times for 1 and 25 min. each); (c) methylene chloride; (d) ethanol; (e) methylene chloride; and (f) 10% triethylamine in chloroform.

The neutralized resin is stirred with alpha-t-butoxycarbonyl(Boc)-leucine 'and 15 diisopropylcarbodiimide (1.5 mmole each) in methylene chloride for 1 hour, and the resulting amino acid resin is then cycled through steps (a) to (f) in the above wash program. Boc-leucine aldehyde (1.25 mmoles), prepared by the method of Fehrentz and Castro, 20 Synthesis, p. 676 (1983), is dissolved in 5 ml of dry dimethylformamide (DMF) and added to the resin TFA salt suspension followed by the addition of 100 mg (2 mmoles) of sodium cyanoborohydride (Sasaki and Coy, Peptides 8:119-121 (1987); Coy et al., id.). After stirring for 25 1 hour, the resin mixture is found to be negative to ninhydrin reaction (1 min.), indicating complete derivatization of the free amino group.

The following amino acids (1.5 mmole) are then coupled successively in the presence disopropylcarbodiimide (1.5 mmole), and the resulting amino acid resin is cycled through washing/deblocking steps (a) to (f) in the same procedure as above:

Boc-His(benzyloxycarbonyl), Boc-Gly, Boc-Val, Boc-Ala,

Boc-Trp, Boc-Gln (coupled in the presence of equivalent of hydroxybenzotriazole), Boc-Asn (coupled in the presence of 1 equivalent of hydroxybenzotriazole), Boc-Gly (coupled as a 6 M excess of the p-nitrophenyl ester), Boc-Leu, Boc-Arg(tosyl), Boc-Gln (coupled as a 6 M excess of the p-nitrophenylester), and pGlu. The completed resin is then washed with methanol and air dried.

The resin described above (1.6 g, 0.5 mmole) is mixed with anisole (5 ml) and anhydrous hydrogen 10 fluoride (35 ml) at 0°C and stirred for 45 min. Excess hydrogen fluoride is evaporated rapidly under a stream of dry nitrogen, and free peptide is precipitated and washed with ether. The crude peptide is dissolved in a minimum volume of 2 M acetic acid and eluted on a column 15 (2.5 x 100 mm) of Sephadex G-25 (Pharmacia Fine Chemicals, Inc.). Fractions containing a major component by uv absorption and thin layer chromatography (TLC) are then pooled, evaporated to a small volume and applied to a column (2.5 x 50 cm) of 20 octadecylsilane-silica (Whatman LRP-1, 15-20 μm mesh size).

The peptide is eluted with a linear gradient of 0-30% acetonitrile in 0.1% trifluoroacetic acid in

25 water. Fractions are examined by TLC and analytical high performance liquid chromatography (HPLC) and pooled to give maximum purity. Repeated lyophilization of the solution from water gives 60 mg of the product as a white, fluffy powder.

The product is found to be homogeneous by HPLC and TLC. Amino acid analysis of an acid hydrolysate confirms the composition of the peptide. The presence of the Leuw[CH2-NH]Leu bond is demonstrated by fast atom bombardment mass spectrometry.

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pGlu-Gln-Arg-Leu-Gly-Asn-Gln-Trp-Ala ψ [CH2-NH]Val-Gly-His-Leu-Met-NH2 and pGlu-Gln-Arg-Leu-Gly-Asn-Gln-Trp-Ala-Val-Gly-His-Leu ψ [CH2NH]Met-NH2 are prepared in similar yields in an analogous fashion by appropriately modifying the above procedure.

A statine or AHPPA residue can be substituted in place of any two amino acids of the peptide, where the peptide contains no pseudopeptide bonds. For example, sta¹³-des Met¹⁴ bombesin was prepared in an analagous fashion by first coupling statine to the resin and then proceeding with the addition of Boc-His(benzylocarbonyl). Statine or Boc-statine can be synthesized according to the method of Rich et al., 1978, J. Organic Chem. 43; 3624; and Rich et al., 1980,

J. Med. Chem. 23: 27, and AHPPA can be synthesized according to the method of Hui et al., 1987, J. Med. Chem. 30: 1287.
Synthesis of Sta 13 - Des - Met 14 Bombesin

solid-phase synthesis of the peptide

pGlu-Gln-Arg-Leu-Gly-Asn-Gln-Trp-Ala-Val-Gly-His-Sta-NH2 was accomplished through the use of the following procedures in which alpha-t-butoxycarbonyl statine (prepared by the procedure of Rich et al., J. Org. Chem. 1978, 43, 3624) is first coupled to

25 methylbenzhydrylamine-polystyrene resin. After
 acetylation, the intermediate
 p-Glu-Gln-Arg(tosyl)-Leu-Gly-Asn-Gln-Trp-Ala-Val-Gly His(benzyloxycarbonyl)-Sta-methylbenzhydrylamine resin
 is prepared. The synthetic procedure used for this
30 preparation follows in detail:

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 Incorporation of alpha-t-butoxycarbonyl statine on methylbenzhydrylamine resin.

Methylbenzhydrylamine-polystyrene resin (Vega Biochemicals, Inc.) (1.0 g, 0.73 mmol) in the chloride ion form is placed in the reaction vessel of a Vega 250C Coupler peptide synthesizer. The synthesizer was programmed to perform the following reactions: (a) methylene chloride; (b) 10% triethylamine in chloroform; (c) methylene chloride; and (d) dimethylformamide.

The neutralized resin is mixed for 18 hours with the preformed active ester made from alpha-t-butoxycarbonyl statine (1.46 mmol), diisopropyl carbodiimide (2 mmol), and hydroxybenzotriazole hydrate (1.46 mmol in dimethylformamide at 0° C. for one hour.

The resulting amino acid resin is washed on the synthesizer with dimethylformamide and then methylene chloride. The resin mixture at this point was found by the Kaiser ninhydrin test (5 minutes) to have an 84% level of statine incorporation on the resin.

Acetylation was performed by mixing the amino-acid resin for 15 minutes with N-acetyl imidazole (5 mmol) in methylene chloride. Derivitization to the 94-99% level of the free amino groups of the resin was indicated by the Kaiser ninhydrin test (5 minutes). The Boc-statine-resin is then washed with methylene chloride.

2. Couplings of the Remaining Amino Acids.

The peptide synthesizer is programmed to perform the following reaction cycle: (a) methylene chloride; (b) 33% trifluroacetic acid (TFA) in methylene chloride (2 times for 5 and 25 min. each); (c) methylene chloride; (d) isopropyl alcohol; (e) 10% triethylamine in chloroform; and (f) methylene chloride.

The following amino acids (2.19 mmol) are then coupled successively by diisopropyl carbodiimide (4

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mmol) alone or diisopropyl carbodiimide (4 mmol) plus hydroxybenzotriazole hydrate (1.47 or 0.73 mmol) and the resulting peptide-resin is washed on the synthesizer with dimethylformamide and then methylene chloride, and then cycled through the washing and deblocking steps (a) to (f) in the procedure described above.

Boc-His (benzyloxycarbonyl) (coupled in the presence of 2 equivalents hydroxybenzotriazole); Boc-Gly; Boc-Val; Boc-Ala; Boc-Trp; Boc-Gln and Boc Asn (coupled as the preformed hydroxybenzotriazole active esters made by reaction at 0° C. for one hour with 1 equivalent hydroxybenzotriazole hydrate); Boc-Gly; Boc-Leu; Boc-Arg(tosyl), Boc-Gln, and pGlu (also coupled as the preformed active esters of hydroxybenzotriazole made by reaction at 0° C. for one hour with 1 equivalent 15 hydroxybenzotriazole hydrate). The completed peptide-resin is then washed with methanol and air dried.

The peptide-resin described above (1.60 g, 0.73 mmol) is mixed with anisole (2.5 mL), dithiothreitol (50 mg), and anhydrous hydrogen fluoride (30 mL) at 0° C. for one hour. Excess hydrogen fluoride is evaporated rapidly under a stream of dry nitrogen, and the free peptide is precipitated and washed with ether. crude peptide is dissolved in 100 mL of 1 M acetic acid and the solution is then evaporated under reduced The crude peptide is dissolved in a minimum volume of methanol/water 1/1 and triturated with 10 volumes of ethyl acetate.

The triturated peptide is applied to a column (9.4 mm I.D. \times 50 cm) of octadecylsilane-silica (Whatman 30 Partisil 10 ODS-2 M 9). The peptide is eluted with a linear gradient of 20-80% of 20/80 0.1% trifluoroacetic acid/acetonitrile in 0.1% trifluoroacetic acid in water. Fractions are examined by TLC and analytical

high performance liquid chromatography (HPLC) and pooled to give maximum purity. Lyophilization of the solution from water gives 77 mg of the product as a white fluffy powder.

Other compounds can be prepared as above and tested for effectiveness as agonists or antagonists in the following test program.

<u>Phase 1 - 3T3 Peptide Stimulated [3H] Thymidine</u> <u>Uptake Assay</u>

10 Cell Culture. Stock cultures of Swiss 3T3
cells (American Type Culture Collection No. CCL 92) are
grown in Dulbecco's Modified Eagles Medium (DMEM)
supplemented with 10% fetal calf serum in humidified
atmosphere of 10% CO₂/90% air at 37°C. For
15 experimental use, the cells are seeded into 24-well
cluster trays and used four days after the last change
of medium. The cells are arrested in the G1/G0 phase of
the cell cycle by changing to serum-free DMEM 24 hours
prior to the thymidine uptake assay.

Assay of DNA Synthesis. The cells are washed 20 twice with 1ml aliquots of DMEM (-serum) then incubated with DMEM (-serum), $0.5\mu M$ [methyl- 3H] thymidine (20Ci/mmole, New England Nuclear), bombesin (1nM), and four concentrations of the test compounds (1, 10, 100, 1000nM) in a final volume of 0.5ml. After 28 hours at 25 37° C, [methyl- 3 H] thymidine incorporation into acid-insoluble pools is assayed as follows. The cells are washed twice with ice-cold 0.9% NaCl (lml aliquots), and acid soluble radioactivity is removed by a 30 min. (4°C) incubation with 5% trichloroacetic acid (TCA). 30 The cultures are then washed once (1ml) with 95% ethanol and solubilized by a 30 min. incubation (lml) with 0.1N The solubilized material is transferred to vials containing 15ml ScintA (Packard), and the radioactivity

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is determined by liquid scintillation spectrometry.

Phase 2 - Small Cell Carcinoma (SCLC) - Bombesin

Stimulated [3H] Thymidine Uptake Assay

Cell Culture. Cultures of the human cell carcinoma cell line (NCI-H69) (obtained from the American Type Culture Association) are maintained in RPMI 1640 medium supplemented with 10% fetal calf serum in 10% CO₂/90% air at 37°C. Twenty-four hours prior to assay, the cells are washed with serum-free medium and seeded in 24-well cluster trays.

Assay of DNA Synthesis. Bombesin (lnM), 0.5µM [methyl-3H] thymidine (20 Ci/mmole, New England Nuclear), and four concentrations of the test compounds (1, 10, 100, 1000nM) are added to the cultures to achieve a final volume of 0.5 ml. After a 28 hr incubation at 37°C, the cells are collected onto GF/B glass fiber filters, and the DNA is precipitated with ice-cold TCA. [3H] thymidine incorporation into acid-insoluble fractions of DNA is determined by liquid scintillation spectrometry.

Phase 3 - Peptide-Induced Pancreatitis

Male, Sprague-Dawley rats (250g) are used for these experiments. The test compound, or 0.9% NaCl is administered s.c. 15 min. prior to the bombesin injection. Bombesin injections are given s.c. at a dose of 10 μg/kg, and blood samples are obtained at 1 hr.30 min., 3hr. and 6hr. Plasma amylase concentration are determined by the Pantrak Amylase test.

Phase 4- In Vitro Inhibition of [125] Gastrin

Releasing Peptide (GRP) Binding to Bombesin

Receptors

Membranes from various tissues (rat brain, rat pancreas, rat anterior pituitary, SCLC, 3T3 cells) are prepared by homogenization in 50mM TrisHCl containing

0.1% bovine serum albumin and 0.1mg/ml bacitracin followed by two centrifugations (39,000xgx15 min., 4°C) with an intermediate resuspension in fresh buffer. For assay, aliquots (0.8ml) are incubated with 0.5nM $[^{125}I]$ GRP ('2000 Ci/mmol, Amersham Corp.) and various 5 concentrations of the test compounds in a final volume of 0.5ml. After a 30 minute incubation at 4°C, the binding reaction is terminated by rapid filtration through Whatman GF/C filters that have been pre-soaked in 0.3% aqueous polethyleneimine to reduce the level of 10 nonspecific binding. The filters and tubes are washed three times with 4ml aliquots of ice-cold buffer, and the radioactivity trapped on the filters is counted by gamma-spectrometry. Specific binding is defined as the total [125] GRP bound minus that bound in the presence 15 of 1000nM bombesin.

Phase 5- Inhibition of Gastrin Release

The stomachs of anesthetized rats are perfused with saline collected over 15 minute periods via pyloric cannulation while the test peptide is infused through the femoral vein for periods between 0 and 150 minutes. Results of Tests of Test Peptides

A number of analogs of bombesin, each containing a non-peptide bond, were synthesized and tested in one or more of the above-described Phase I - 5 assays; the results of Phase 1, 2, and 4 tests are given in Table 1 attached hereto (analogs of bombesin are indicated by the symbol "BN"). Brain and 3T3 GRP receptor and thymidine uptake data are expressed in IC50 (nM). Table 1 also gives results for non-peptide bond-containing analogs of three other naturally-occurring peptides, Substance P (which plays a role in the sensation of pain), Neuromedin C, whose C-terminal seven amino acids are similar to those of

bombesin, and litorin, whose eight C-terminal amino acids are identical to Bombesin, with the exception of a Phe substitution for Leu at position A¹³ of bombesin.

In Table 1, the position of the non-peptide

5 bond is indicated by the position of the symbol \(\psi \);

i.e., \(\psi \) is always shown preceding the amino acid

which, in that peptide, is bonded to the amino acid

N-terminal to it via the non-peptide bond. Where no

amino acid is specified under "structure", as in

BIM-26034, the non-peptide bond links the two peptides

represented by the numbers given as post-scripts (e.g.,

between amino acids 7 and 8 of BIM-26034, which

otherwise is identical to naturally occurring

bombesin).

In Table 1, it can be seen that a preferred 15 placement of the non-peptide bond in bombesin analogs is at the 13-14 position; two of the most active analogs (as indicated by a low GRP receptor IC50 value) are BIM-26027 and BIM-26028. However, BIM-26027 causes proliferation of cancer cells (see Table 1, under 20 thymidine uptake), and therefore is an agonist and not an antagonist. In general, compounds having the non-peptide bond at any position other than the active site of the peptide are agonists rather than antagonists. Table I also shows that when statine 25 replaces the ${\rm A}^{13}$ and ${\rm A}^{14}$ residues of bombesin, the resultant bombesin analog BIM-26096 causes proliferation of cancer cells and is therefore an agonist. Bombesin superagonists may be useful in cancer therapy, as suggested by Alexander et al., 1988, Cancer Research 48: 30 1439-1441, and Alexander et al., 1988, Pancreas 3:297-302, hereby incorporated by reference. Alexander et al. show that chronic bombesin treatment inhibited the growth of human ductal adenocarcinoma transplanted

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into athymic mice. These results were surprising for bombesin stimulates growth of normal pancreas tissue. The demonstration of both stimulatory and inhibitory activity suggests that bombesin interacts differently in normal and neoplastic pancreatic tissues.

These observations prompted us to evaluate the affect of BIM-26096, a bombesin analogue which has bombesin-like agonist activity, on the <u>in vitro</u> growth of a pancreatic tumor cell line (AR42J). For these experiments, AR42J cells were subcultured into a 24-well culture plate in Dulbecco's modified Eagle's medium containing 10% fetal calf serum containing various concentrations (0.1-100nM) of BIM-26096. After a 36 hr incubation the cells were removed with a trypsin/EDTA solution and the number of cells were determined using a Coulter Counter. The results are shown below:

	Treatment	<pre>Cell Count (% Control)</pre>
	control	100
	BIM-26096 (0.1 nM)	78
20	BIM-26096 (1.0 nM)	73
	BIM-26096 (10 nM)	56
	BIM-26096 (100 nM)	52

These results indicate that the bombesin agonist, BIM-26096, has <u>in vitro</u> antiproliferative activity against the AR42J rat pancreas tumor.

Finally, Table 1 also shows that bond placement, while important, is not the only factor influencing antagonist activity, and that amino acid substitutions at some positions exert influence as well; this is illustrated by BIM-26030, with Gly in position 11, which exhibited no antagonist activity. Table 1 also gives negative results for analogs of Spantide ([D-Arg', D-Trp^{7,9}, Leu"] Substance P, and Bombesin. Thus the non-peptide bond placement quidelines given

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herein should be used in conjunction with the routine assays described above to select useful antagonists or agonists.

In a phase 5 assay, above, the results of which are not given in Table 1, BIM-26028 was shown to be a potent inhibitor of bombesin - stimulated gastric acid secretion.

Use

The peptides of the invention may be

administered to a mammal, particularly a human, in one
of the traditional modes (e.g., orally, parenterally,
transdermally, or transmucosally), in a sustained
release formulation using a biodegradable biocompatible
polymer, or by on-site delivery (e.g., in the case of
anti-cancer bombesin to the lungs) using micelles, gels
and liposomes.

The bombesin antagonists and agonists of the invention are suitable for the treatment of all forms of cancer where bombesin-related substances act as autocrine or paracrine mitotic agents, particularly small-cell lung carcinoma. The peptides can also be used for the inhibition of gastric acid secretion, the symptomatic relief and/or treatment of exocrine pancreatic adenocarcinoma, and the restoration of appetite to cachexic patients. The peptides can be administered to a human patient in a dosage of 0.5 µg/kg/day to 5 mg/kg/day. For some forms of cancer, e.g., small cell lung carcinoma, the preferred dosage for curative treatment is 250mg/patient/day.

Other Embodiments

Other embodiments are within the following claims.

For example, as is mentioned above, there are a number of other peptide families from which agonists or 5 antagonists can be made according to the invention. Some of these families are substance P and related peptides, vasoactive inestial peptide (VIP) and related peptides, and neurotensin and related peptides. number of peptides in each family on which antagonists 10 or agonists can be based is large. For example, there are at least 10 currently-known peptides in the VIP family, including sauvagine and urotensin. In addition, there have been isolated seven natural bradykinin-like peptides. Neurotensin 15 (pGlu-Leu-Tyr-Glu-Asn-Lys-Pro-Arg-Arg-Pro-Tyr-Ile-Leu-OH) has two peptide bonds which advantageously can be replaced by non-peptide bonds: Ile-Leu and Tyr-Ile. In addition, neurotensin antagonists can be missing any or all of the N-terminal 20 seven amino acid residues, as it has been shown (Granier et al. (1984) Eur. J. Biochem. 124: 117) that they are not needed for biological activity and binding. Screening of neurotensin antagonists can be by binding 25 to SCLC receptors. Gastrin releasing peptides (GRP) and related peptides (e.g., Neuromedin C (GRP 18-27)) have a bond between amino acid residues 13 and 14 which can be replaced with a non-peptide bond to form a GRP antagonist.

- · 23 -

Table 1

<u>Code</u>	<u>Structure</u>	Brain GRP Receptor <u>IC50(nM)</u>	3T3 GRP Receptor IC50(nM)	Thym. Uptake IC50(nM)
BIM-26025	[His^{12} Y[CH_2 NH] Leu^{14}]BN	>1000		
BIM-26026	$[Ala^9 \Psi [CH_2 NH] Leu^{14}] BN$	>1000		1574
BIM-26027	[Val ¹⁰ Y[CH ₂ NH]Leu ¹⁴]BN	0.48	2.3	agonsit EC50=0.07n
M BIM-26028	$[Leu^{13}Y[CH_2NH]Leu^{14}]BN$	13		14.7
BIM-26030	$[Gly^{11}Y[CH_2NH]Leu^{14}]BN$	>1000		• :
BIM-26034	[Y[CH2NH]8,7]BN	>1000		:
BIM-26036	[Des-pGlu ¹ ,Gln ² ,Y(Ala ⁹ , Val ¹⁰)Phe ¹²]BN	>1000		
BIM-26046	$[Gly^{11}Y[CH_2NH]D-Phe^{12},$ Leu ¹⁴]BN	>1000		:
BIM-26048	$[D-Phe^{12}Y[Ch_2NH]Leu^{13},$ $Leu^{14}]BN$	>1000		
BIM-26056	[Leu ¹⁰ Y[CH ₂ NH] Leu ¹¹ NH ₂]Substance P	>1000		:
BIM-26057	(Cys ⁹ , wLeu ¹³ , Cys ¹⁴)BN	>1000	•	:

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<u>Cođe</u>	Structure	Brain GRP Receptor IC50(nM)	_	Thym. Uptake
		1C30(IIII)	IC50(nM)	IC50(nM)
BIM-26061	[D-pGlu,D-Ala ⁵ ,YLeu ⁷ , Met ⁸]BN	>1000		
BIM-26062	[YPhe ¹³ ,Leu ¹⁴]BN	>1000		437
BIM-26063	[des-Glu ⁷ , YLeu ¹³⁻¹⁴]BN	>1000		
BIM-26064	[ψ Leu ¹⁰ ,Nle ¹¹]Spantide	>1000		
BIM-26067	$[\operatorname{des-Gln}^7, \operatorname{\psi Leu}^{13-14}]$ BN	>1000	·	
BIM-26068	$[\psi Leu^{13}, Phe^{14}]$ BN	2.9		70
BIM-26070	[\psi D-Trp9,Nle11]Spantide	>1000		
BIM-26071	[Tyr ⁴ , wLeu ¹³ [CH ₂ NH]-Met ¹⁴]BN	34	16	104
BIM-26072	[Cys ⁹ ,Leu ¹³ [CH ₂ NH] Cys ¹⁴]BN	>1000		
BIM-26073	[Cys ⁹ , wLeu ¹³ [CH ₂ NH] Cys ¹⁴]BN	>1000		
BIM-26074	[Des-Gln ⁷ , \psi Leu ¹³ [CH ₂ NH] Leu ¹⁴]BN	>1000		
BIM-26075	[D-Phe ¹¹ , wLeu ¹³⁻¹⁴]BN	>1000		

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<u>Code</u>	Structure	Brain GRP Receptor IC50(nM)	3T3 GRP Receptor IC50(nM)	Uptake
BIM-26076	[D-Phe ¹¹ , \psi Leu ¹³⁻¹⁴]BN	>1000		
BIM-26077	[D-Ala ⁵ , wLeu ¹³⁻¹⁴]BN	517	196	1001
BIM-26078	[D-Ala ¹¹ , wLeu ¹³⁻¹⁴]BN	>1000		70
BIM-26079	[\psiPhe 7, Leu 11] Spantide	>1000		
BIM-26080	[\psi Gln 6-Nle 11]Spantide	>1000		
BIM-26081	[\psi D-Trp 7-Nle 11]Spantide	>1000		
BIM-26082	[\psiPhe 8-Nle 11]Spantide	>1000		
BIM-26083	[\psi GLn 6-Nle 11] Spantide	>1000		
BIM-26084	[\psi D-Trp 7-Nle 11] Spantide	>1000		
`BIM-26085	[\psi Phe 8-N1e 11] Spantide	>1000		•
BIM-26086	[D-Phe ¹² , wLeu[CH ₂ NH] Leu ¹⁴]EN	>1000		
BIM-26088	[\psi Gly 9 [CH2NH] Leu 14] Spantide	>1000		
BIM-26089	[\psi Gln 6 [CH 2NH] Leu 11] Spantide	>1000		
BIM-26090	[YPhe ⁷ ,Leu ¹¹]Substance P			>1000

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		Brain GRP	3T3 GRP	•
Code	Structure	Receptor <u>IC50(nM)</u>	Receptor IC50(nM)	
BIM-26091	[ψPhe ⁸ ,Leu ¹¹]Substance P			>1000
BIM-26092	[\psi Leu 9] Neuromedin C		. 24	466
BIM-26093	[D-Ala ¹ ,ψ[CH ₂ NH]Leu ⁹] Neuromedin C			2 171
BIM-26094	$[D-Ala^{5,11}, Leu^{13}\psi[CH_2NH]]$ $Leu^{14}]BN$		161	3 574
BIM-26095	[D-Ala ⁶ , Leu ⁹ ψ[CH ₂ NH] Leu ¹⁰]Litorin		262	3 1209
BIM-26096	[Sta ¹³ ,Des Met ¹⁴]BN	33		agonsit EC50=3nM
BIM-26097	[Ac-Lys ⁷ , wLeu ¹³]BN ₇₋₁₄	1000		>1000
BIM-26098	[Lys ⁷ , \psi Leu ¹³] BN ₇₋₁₄	1000	•	
BIM-26099	[\psi Leu 13, Met]BN		73	78
BIM-26100	[Phe ⁸ ψ[CH ₂ NH]Leu ⁹]Litorin		74	22
BIM-26101	Leu ⁸ ψ[CH ₂ NH]Leu ⁹]Litorin		17.9	257
BIM-26102	ψPhe ⁹ [CH ₂ NH]Met ¹⁰ NH ₂ Neuromedin β		184	>1000

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	•	Brain GRP Receptor	3T3 GRP Receptor	Thym. Uptake
Code	Structure	IC50(nM)	IC50(nM)	IC50(nM)
BIM-26103	ψLeu ¹³ [CH ₂ NH]Met ¹⁴ NH2 A-Lytensin		>1000	>1000
BIM-26104	ψLeu ⁷ [CH ₂ NH]Met ⁸ NH ₂ GRP(20-27)			>1000
Spantide	[D-Arg ¹ ,D-Trp ^{7,9} ,Leu ¹¹] Substance P		3303	2171
Bombesin	pGlu-Gln-Arg-Leu-Gly-Asn- Gin-Trp-Ala-Val-Gly-His- Leu-Met-NH ₂	15	0.17	

Claims

- A linear peptide which is an analog of naturally occurring, biologically active bombesin having an active site and a binding site responsible for the binding of bombesin to a receptor on a target cell, cleavage of a peptide bond in said active site of said naturally occurring bombesin being unnecessary for in vivo biological activity of bombesin, said analog having a non-peptide bond instead of a peptide bond between an amino acid of said active site and an adjacent amino 10 acid, said analog being capable of binding to said receptor, so that said analog is capable of acting as a competitive inhibitor of said naturally occurring peptide by binding to said receptor and, by virtue of said non-peptide bond, failing to exhibit the in vivo 15 activity of said naturally occurring bombesin.
- 2. The linear peptide of claim 1 wherein said naturally occurring bombesin is characterized in that one or more amino acids in the amino terminal half of bombesin are hydrogen bonded to one or more amino acids in the carboxy terminal half of bombesin, and said non-peptide bond of said linear peptide decreases said hydrogen bonding.
- 3. The linear peptide of claim 2 wherein said hydrogen bonded amino acids of said naturally occurring bombesin make up at least a portion of the active site of said naturally occurring bombesin, so that said active site is inactivated by the decrease in hydrogen bonding caused by said non-peptide bond.

- 4. A linear peptide which is an analog of naturally occurring, biologically active human bombesin which includes an active site comprising at least one amino acid in the carboxy terminal half of bombesin, said linear peptide including said amino acid in its carboxy terminal half, there being a non-peptide bond bonding said amino acid to an adjacent amino acid.
- 5. The linear peptide of claim 4 wherein said amino acid of said naturally occurring bombesin is hydrogen bonded to another, non-adjacent amino acid in said bombesin, and said non-peptide bond in said linear peptide causes a decrease in said hydrogen bonding which inactivates said bombesin.
- 6. An effective bombesin antagonistic peptide containing the amino acid formula:

wherein

 $A^1 = pGlu \text{ or is deleted};$

20 $A^2 = Gln$, Asn, Gly, Ala, Leu, Ile, Nle, α -aminobutyric acid, Met, Val, Phe, p-X-Phe (X = F, Cl, Br, OH or CH₃), Trp, β -naphthylalanine or is deleted;

 A^3 = Arg, D-Arg, Lys, D-Lys or is deleted; A^4 = Gln, Asn, Gly, Ala, Leu, Ile, Nle, α -aminobutyric acid, Met, Val, Phe, p-X-Phe (X = F, Cl, Br, OH or CH₃), Trp, β -naphthylalanine or is deleted;

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A^5 =
                Gln, Asn, Gly, Ala, Leu, Ile, Nle,
                \alpha-aminobutyric acid, Met, Val, Phe, D-Phe,
                p-X-Phe (X = F, Cl, Br, OH or CH<sub>3</sub>), Trp,
                B-naphthylalanine, D-Ala or is deleted;
      A^6 =
5
                Gln, Asn, Gly, Ala, D-Ala, N-Ac-D-Ala, Leu,
                Ile, Nle, \alpha-aminobutyric acid, Met, Val, Phe,
                p-X-Phe (X = F, Cl, Br, OH or CH<sub>3</sub>), Trp,
                p-Glu, ß-naphthylalanine or is deleted;
                Gln, Asn, Gly, Ala, Leu, Ile, Nle,
                lpha-aminobutyric acid, Met, Val, Phe, D-Phe,
10
                p-X-Phe (X = F, Cl, Br, OH or CH<sub>3</sub>), Trp, His,
                or B-naphthylalanine;
                Trp;
               Gln, Asn, Gly, Ala, Leu, Ile, Nle,
15
               \alpha-aminobutyric acid, Met, Val, Phe, p-X-Phe
               (X = F, Cl, Br, OH or CH<sub>3</sub>), Trp, or
               B-naphthylalanine;
               Gln, Asn, Gly, Ala, Leu, Ile, Nle,
               lpha-aminobutyric acid, Met, Val, Phe, p-X-Phe
20
               (X = F, C1, Br, OH or CH<sub>3</sub>), Trp, or
               ß-naphthylalanine;
    A<sup>11</sup>
               Gly, or D-Ala;
    A^{12} =
               His, Phe, or p-X-Phe (X = F, Cl, Br, OH, CH_3);
    A^{13} =
               Gln, Asn, Gly, Ala, Leu, Ile, Nle,
25
               \alpha-aminobutyric acid, Met, Val, Phe, p-X-Phe
               (X = F, Cl, Br, OH or CH<sub>3</sub>), Trp, or
               B-naphthylalanine;
              Gln, Asn, Gly, Ala, Leu, Ile, Nle,
              \alpha-aminobutyric acid, Met, Val, Phe, p-X-Phe
30
              (X = F, Cl, Br, OH or CH<sub>3</sub>), Trp, or
              ß-naphthylalanine;
    provided that
              each R_1, R_2, R_3, and R_4, independently,
    is H, C_{1-12} alkyl, C_{7-10} phenylalkyl, COE_1 (where
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 E_1 is C_{1-20} alkyl, C_{3-20} alkenyl, C_{3-20} alkinyl, phenyl, naphthyl, or C₇₋₁₀ phenylalkyl), or COOE₂ (where E_2 is C_{1-10} alkyl or C_{7-10} phenylalkyl), and R_1 and R_2 are bonded to the N-terminal amino acid of said peptide, which can be A¹, A², A³, A^4 , A^5 , A^6 , or A^7 , and further provided that when one of R_1 or R_2 is COE_1 or $COOE_2$, the other must be H, and when one of R3 or R4 is COE1 or COOE, the other must be H, and further provided that when $A^1 = pGlu$, R_1 must be H and R_2 must be the 10 portion of Glu that forms the imine ring in pGlu; and for each of the residues A^7 , A^8 , A^9 , A^{10} , A^{11} , A^{12} , and A^{13} , independently, the carbon atom participating in the amide bond between that residue and the nitrogen atom of the alpha amino group of the adjacent amino acid residue may be a carbonyl carbon or may be reduced to a methylene carbon, provided that at least one such carbon atom must be reduced to a methylene carbon; or a pharmaceutically acceptable salt thereof. 20

- 7. The effective bombesin antagonist peptide of claim 6 wherein A^1 through A^6 are deleted and the carbon atom participating in the amide bond between Leu¹³ and Leu¹⁴ is a methylene carbon; or a pharmaceutically acceptable salt thereof.
- 8. The effective bombesin antagonist peptide of claim 6 wherein, for each of said residues A¹¹, A¹², and A¹³, independently, the carbon atom participating in the amide bond between that residue and the nitrogen atom of the alpha amino group of the adjacent amino acid residue may be a carbonyl carbon or may be reduced to a methylene carbon, provided that at least one such carbon atom must be reduced to a

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methylene carbon; or a pharmaceutically acceptable salt thereof.

9. An effective litorin antagonist peptide containing the amino acid formula:

Ala; A⁵ is Val; A⁶ is Gly or D-Ala; A⁷ is His; A⁸ is Phe or Leu; and A⁹ is Met or Leu; provided that the carbon atom participating in the amide bond between the A⁸ residue and the nitrogen atom of the alpha amino group of the adjacent amino acid residue may be a carbonyl carbon or may be reduced to a methylene carbon; or a pharmaceutically acceptable salt thereof.

- 10. An effective bombesin agonist of the
 general formula of claim 6 wherein, for each of the
 residues A⁹, A¹⁰, A¹¹, A¹², A¹³, and A¹⁴,
 independently, the carbon atom participating in the
 amide bond between that residue and the nitrogen atom of
 the alpha amino group of the adjacent amino acid residue
 may be a carbonyl carbon or may be a non-peptide bond,
 provided that said non-peptide bond is said carbonyl
 carbon having been reduced to a methylene carbon,
 further provided that at least one such carbon atom must
 be reduced to a methylene carbon; or a pharmaceutically
 acceptable salt thereof.
 - 11. A bombesin agonist having the amino acid formula

pGlu-Gln-Arg-Leu-Gly-Asn-Gln-Trp-Ala-Val-Gly-His-Leu-Leu[Val 10 Y[CH $_2$ NH]Leu 14]BN.

- 12. An effective bombesin agonist having the amino acid formula of claim 6 which is an analog of naturally occurring, biologically active bombesin having an active site, said active site includes the positions A^9 , A^{10} , A^{11} , A^{12} , A^{13} , and A^{14} , and a 5 binding site responsible for the binding of said bombesin to a receptor on a target cell, said analog having either (a) said non-peptide bond at residues other than within said active site, or (b) having at least one statine or AHPPA residue in place of two 10 naturally occurring amino acids of said active site, and further provided that the peptide can contain statine or AHPPA when all bonds between amino acid residues are peptide bonds, and further provided that when an amino acid residue is statine or AHPPA, the amino acid to the 15 right of it in the formula is deleted, so that said analog is capable of binding to said receptor, and, by virtue of said statine or AHPPA residue, exhibiting enhanced in vivo biological activity compared to said naturally occurring bombesin. 20
 - 13. A bombesin agonist having the amino acid formula

pGlu-Gln-Arg-Leu-Gly-Asn-Gln-Trp-Ala-Val-Gly-His-[Sta¹³, Des Met¹⁴].

INTERNATIONAL SEARCH REPORT

International Application No. PCT/US88/03286 I. CLASSIFICATION OF SUBJECT MATTER (if several classification symbols apply, indicate all) 6 According to International Patent Classification (IPC) or to both National Classification and IPC IPC(4): C07K 7/02, 7/06, 7/08 U.S. CL: 530/327, 328, 323 II. FIELDS SEARCHED Minimum Documentation Searched 7 Classification System Classification Symbols 530/327, 328, 323 U.S. Documentation Searched other than Minimum Documentation to the Extent that such Documents are Included in the Fields Searched 8 Chemical Abstracts and Biological Abstracts Online Computer Search. III. DOCUMENTS CONSIDERED TO BE RELEVANT 9 Category 5 Citation of Document, 11 with indication, where appropriate, of the relevant passages 12 Relevant to Claim No. 13 US, A, 4,207,311 (Brown et. al.), 10, June 1980. See column 2, line 29 in particular. Am J. of Physiol, (Maryland, USA) 1-13 issued 1986, (Heinz-Erian et. al.), "[D-Phe12] bombesin analogues: a new class of bombesin receptor antagonists", pages G439-G442. Proc. Natl. Acad. Sci. USA (Washington, 1-13 D.C., USA) volume 82, issued November, 1985. (Zachary et. al.), "Highaffinity receptors for peptides of the bombesin family in Swiss 3T3 cells", pages 7616-7620. Special categories of cited documents: 10 "T" later document published after the International filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "A" document defining the general state of the art which is not considered to be of particular relevance earlier document but published on or after the international filing date "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled document referring to an oral disclosure, use, exhibition or other means document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family IV. CERTIFICATION Date of the Actual Completion of the International Search Date of Malling of this International Search Report 22 December 1988 International Searching Authority Signature of Authorized Officer Chirostonia Chan ISA/US Christina Chan

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III. DOCUMENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET) Category Citation of Document, 16 with Indication, where appropriate, of the relevant passages 17 Relevant to Claim No 18					
Category *	Citation of Document, 22 Will indication, where appropriate, or the research passages	Relevant to Glass tto			
A	J. Med. Chem. (Washington, D.C., USA) volume 28 issued 1985, (Martinez et. al.), "synthesis and biological activities of some pseudo-peptide analogues of tetragastrin: The importance of the peptide backbone", pages 1874-1879.	1-13			
A	J. Med. Chem. (Washington, D.C., USA) volume 30, issued 1987, (Rodriguez et. al.). "Synthesis and biological activities of Pseudopeptide analogues of the C-terminal heptapeptide of cholecystokinin. On the importance of the peptide bonds", pages 1366-1373.	1-13			
¥	J. Med. Chem. (Washington, D.C. USA) volume, 30, issued 1987, (Sasaki et. al.), "Solid-Phase Synthesis and biological Properties of [CH2NH] Pseudopeptide analogues of a highly potent somatostatin octapeptide", pages 1162-1166. See pages 1162, 1164, 1166 in particular.	1-8 10-13			
Y	Cancer Surveys (Oxford, England) volume 4, No. 4, issued 1985 (Cuttitta et. al.), "Autocrine growth factors in human small cell lung cancer", pages 707-727. See page 718 in particular.	1-8 10-13			
X,P	Chemical Abstract, (Columbus, Ohio, USA) volume 109, issued 1988, (Coy et. al.), "Probing peptide backbone function in bombesin. A reduced peptide bond analog with potent and specific receptor antagonist activity", the abstract No. 32216K, J. Biol. Chem. 1988, 263 (11), 5056-60 (Eng).	1-8 10-13			